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Transmitted herewith for filing under 37 CFR 1.53(b) is the

- ☐ patent application of  
☐ continuation patent application of  
☐ divisional patent application of  
☒ continuation-in-part patent application of

Inventor(s)/Applicant Identifier: Peter S. Lu

For: CLASP-2 TRANSMEMBRANE PROTEINS

☐ This application claims priority from each of the following Application Nos./filing dates:

the disclosure(s) of which is (are) incorporated by reference.

☐ Please amend this application by adding the following before the first sentence: "This application is a ☐ continuation ☐ continuation-in-part of and claims the benefit of U.S. Provisional Application No. 60/\_\_\_\_\_, filed \_\_\_\_\_, the disclosure of which is incorporated by reference."

Enclosed are:

- ☒ 124 page(s) of specification  
☒ 5 page(s) of claims  
☒ 1 page of Abstract  
☒ 114 sheet(s) of ☐ formal ☒ informal drawing(s).  
☐ An assignment of the invention to \_\_\_\_\_  
☒ A ☐ signed ☒ unsigned Declaration & Power of Attorney  
☐ A ☐ signed ☐ unsigned Declaration.  
☐ A Power of Attorney.  
☐ A verified statement to establish small entity status under 37 CFR 1.9 and 37 CFR 1.27 ☐ is enclosed ☐ was filed in the prior application and small entity status is still proper and desired.  
☐ A certified copy of a \_\_\_\_\_ application.  
☐ Information Disclosure Statement under 37 CFR 1.97.  
☐ A petition to extend time to respond in the parent application.  
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**In view of the Unsigned Declaration as filed with this application and pursuant to 37 CFR §1.53(f),  
Applicant requests deferral of the filing fee until submission of the Missing Parts of Application.**

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**PATENT APPLICATION**  
**CLASP-2 TRANSMEMBRANE PROTEINS**

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## CLASP-2 TRANSMEMBRANE PROTEINS

### 0.0 CROSS-REFERENCES TO RELATED APPLICATIONS

This application is a continuation-in-part application of U.S. Patent Application No. 09/547,276 filed April 11, 2000, which claims the benefit of U.S. Provisional Application Nos. 60/182,296 filed February 14, 2000, 60/176,195 filed January 14, 2000, 60/170,453 filed December 13, 1999, 60/162,498 filed October 29, 1999, 60/160,860 filed October 21, 1999, 60/134,118 filed May 14, 1999, 60/134,117 filed May 14, 1999, 60/134,114 filed May 14, 1999, and 60/129,171 filed April 14, 1999, the disclosures of which are incorporated by reference.

### 1.0 FIELD OF THE INVENTION

The present invention relates to molecules expressed in cells of the immune system. In particular, the invention relates to a transmembrane protein that contains certain classical cadherin characteristics.

### 2.0 BACKGROUND OF THE INVENTION

The generation of an immune response against an antigen is carried out by a number of distinct immune cell types that work in concert within the context of a particular antigen. The helper T cell ( $T_H$ ) plays a pivotal role to coordinate two types of antigen-specific immune response; *i.e.*, cellular and humoral immune response. Recognition of antigen by T cells requires the formation of a specialized junction between the T cell and the antigen-presenting cell (APC) called the "immunological synapse" (Dustin, *et al.*, 1998, Cell 94: 667-677). The immune synapse orchestrates recruitment and exclusion of specific proteins from the contact area by an unknown mechanism and is thought to be initiated by T-cell antigen receptor (TCR) recognition of peptides bound to MHC molecules (antigen) (Monk, *et al.* 1998, Nature 395: 82). However, the low affinity of the TCR for antigen as well as limited number of ligands makes it unlikely that TCR: antigen interaction alone is sufficient to drive the formation of the immunological synapse (Matsui *et al.*, 1994, Proc. Natl. Acad. Sci. U.S.A. 91: 12861-12866).

Costimulatory molecules such as CD4, ICAM-1, LFA-1, CD28, CD2 have been proposed to stabilize the cell-cell contact (Dustin, *et al.*, 1999, Science 283: 649).

However, since these molecules are recruited to the synapse after activation they cannot

account for the high specificity and avidity during the early phases of T-cell antigen recognition. Recent work demonstrated that a portion of the T cell surface at the leading edge is specialized to mediate the early phases of synapse formation (Negulescu, *et al.*, 1996, Immunity 4: 421-430). Such a specialization must be a pre-formed structure containing cell surface adhesion proteins (ectodomains) to augment TCR engagement and corresponding cytoplasmic portions (endodomains) to transduce signals and bind cytoskeleton to maintain structural/functional polarity.

The ectodomain of the pre-formed synapse or "immune gateway" was recently discovered and is created in part by CLASP-1 (U.S.S.N. 09/411,328, filed October 1, 1999; PCT/US99/22996). In addition to cadherin motifs, CLASP-1 also contains a CRK-SH3 binding domain, tyrosine phosphorylation sites, and coiled/coil domains suggesting direct interaction with cytoskeleton and regulation by adaptor molecules such as CRK. The *CLASP-1* transcript is present in lymphoid organs and neural tissue, and the protein is expressed by T and B lymphocytes and macrophages in the MOMA-1 subregion of the marginal zone of the spleen, an area known to be important in T: B cell interaction. CLASP-1 staining of individual T and B cells exhibits a preactivation structural polarity, being organized as a "ball" or "cap" structure in B cells, and forming a "ring", "ball" or "cap" structure in T cells. The placement of these structures is adjacent to the microtubule-organizing center ("MTOC"). CLASP-1 antibody staining indicates that CLASP-1 is at the interface of T-B cell conjugates that are fully committed to differentiation. Antibodies to the extracellular domain of CLASP-1 also block T-B cell conjugate formation and T cell activation.

### 3.0 SUMMARY OF THE INVENTION

The present invention relates to a cell surface molecule, a member of a new multigene family designated cadherin-like asymmetry protein(s) ("CLASP(s)"). In particular, it relates to a polynucleotide comprising a coding sequence for CLASP-2, a polynucleotide that selectively hybridizes to the complement of a CLASP-2 coding sequence, expression vectors containing such polynucleotides, genetically-engineered host cells containing such polynucleotides, CLASP-2 polypeptides, CLASP-2 fusion proteins, therapeutic compositions, CLASP-2 domain mutants, antibodies specific for CLASP-2 polypeptides, methods for detecting the expression of CLASP-2, and methods of inhibiting an immune response by interfering with CLASP-2 function. A wide variety of uses are encompassed by the invention, including but not limited to, treatment of autoimmune



diseases and hypersensitivities, prevention of transplantation rejection responses, and augmentation of immune responsiveness in immunodeficiency states.

In one aspect, the invention provides an isolated CLASP-2 polynucleotide that is: (a) a polynucleotide that has the sequence of SEQ ID NO: 1, 3, 5 or 9; (b) a polynucleotide that hybridizes under stringent hybridization conditions to (a) and encodes a polypeptide having the sequence of SEQ ID NO: 2, 4, 6 or 10 or an allelic variant or homologue of a polypeptide having the sequence of SEQ ID NO: 2, 4, 6 or 10; or (c) a polynucleotide that hybridizes under stringent hybridization conditions to (a) and encodes a polypeptide with at least 25 contiguous residues of the polypeptide of SEQ ID NO: 2, 4, 6 or 10; or (d) a polynucleotide that hybridizes under stringent hybridization conditions to (a) and has at least 12 contiguous bases identical to or exactly complementary to SEQ ID NO: 1, 3, 5, or 9. In a related aspect, the invention provides a CLASP-2 polynucleotide wherein the polynucleotide encodes a polypeptide that binds to the PDZ domain of PSD95, DLG1 or neDLG. 2. In another related aspect, the invention provides a CLASP-2 polynucleotide wherein the polynucleotide encodes a polypeptide that has a binding affinity of at least  $10^4 \text{ M}^{-1}$  for binding PSD95, DLG1 or neDLG.

In one aspect, the invention provides a CLASP-2 polynucleotide that encodes a polypeptide having the full-length sequence of SEQ ID NO: 2, 4, 6, or 10 or the cDNA sequence encoded by the inserts of ATCC Deposit Nos: PTA-1562, PTA-1563 and PTA-1573.

In another aspect, the invention provides a CLASP-2 polynucleotide that encodes a polypeptide having the full-length sequence of Isoform 1, Isoform 2, or Isoform 3 (SEQ ID NO: \_\_\_\_\_) or the cDNA sequence encoded by the inserts of AVC-PD14 (ATCC Deposit No. \_\_\_\_\_) and AVC-PD19 (ATCC Deposit No. \_\_\_\_\_).

In another aspect, the invention further provides an isolated CLASP-2 polynucleotide comprising a nucleotide sequence that has at least 90% percent identity to SEQ ID NO: 1, 3, 5 or 9 as calculated using FASTA wherein said sequences are aligned so that highest order match between said sequences is obtained.

The invention further provides an isolated polypeptide comprising a nucleotide sequence that has at least 90% sequence identity to SEQ ID NO: 2, 4, 6 or 10 and is immunologically crossreactive with SEQ ID NO: 2, 4, 6 or 10 or shares a biological function with native CLASP-2.

The invention also provides vectors, such as expression vectors, comprising a polynucleotide sequence of the invention. In other embodiments, the invention provides host

cells or progeny of the host cells comprising a vector of the invention. In certain embodiments, the host cell is a eukaryote. In other embodiments, the expression vector comprises a CLASP-2 polynucleotide in which the nucleotide sequence of the polynucleotide is operatively linked with a regulatory sequence that controls expression of the polynucleotide in a host cell. In certain embodiments, the invention provides a host cell comprising a CLASP-2 polynucleotide, wherein the nucleotide sequence of the polynucleotide is operatively linked with a regulatory sequence that controls expression of the polynucleotide in a host cell, or progeny of the cell.

In another aspect, the invention further provides a CLASP-2 polynucleotide that is an antisense polynucleotide. In a preferred embodiment, the antisense polynucleotide is less than about 200 bases in length. In other embodiments, the invention provides an antisense oligonucleotide complementary to a messenger RNA comprising SEQ ID NO: 1, 3, 5 or 9 and encoding CLASP-2, wherein the oligonucleotide inhibits the expression of CLASP-2.

In another aspect, the invention provides an isolated DNA that encodes a CLASP-2 protein as shown in SEQ ID NO: 2, 4, 6 or 10. In certain embodiments, the CLASP-2 polynucleotide is RNA.

The invention provides a method for producing a polypeptide comprising: (a) culturing the host cell containing a CLASP-2 polynucleotide under conditions such that the polypeptide is expressed; and (b) recovering the polypeptide from the cultured host cell or its cultured medium.

The invention further provides an isolated CLASP-2 polypeptide encoded by a CLASP-2 polynucleotide. In some embodiments, the CLASP-2 polypeptide has the amino acid sequence of SEQ ID NO: 2, 4, 6 or 10, or a fragment thereof. In some embodiments, the isolated CLASP-2 polypeptide is cell-membrane associated. In other embodiments, the isolated CLASP-2 polypeptide is soluble. In other embodiments, the soluble CLASP-2 polypeptide is fused with a heterologous polypeptide.

The invention further provides an isolated CLASP-2 protein having the sequence as shown in SEQ ID NO: 2, 4, 6 or 10. In some embodiments, the invention provides a CLASP-2 protein comprising the sequence as shown in SEQ. ID. NO: 1 and variants thereof that are at least 95% identical to SEQ ID. NO: 2 and specifically binds a cytoskeletal protein. In certain embodiments the cytoskeletal protein is spectrin.

The invention further provides an isolated antibody that specifically binds to a polypeptide having the amino acid sequence as shown in SEQ ID NO: 2, 4, 6 or 10, or a

binding fragment thereof. In some embodiments the antibody is monoclonal. In other embodiments, the invention provides a hybridoma capable of secreting the antibody.

The invention further provides a method of identifying a compound or agent that binds a CLASP-2 polypeptide comprising: i) contacting a CLASP-2 polypeptide with the compound or agent under conditions which allow binding of the compound to the CLASP-2 polypeptide to form a complex and ii) detecting the presence of the complex.

The invention further provides a method of detecting a CLASP-2 polypeptide in a sample, comprising: (a) contacting the sample with a CLASP-2 antibody or binding fragment and (b) determining whether a complex has been formed between the antibody and with CLASP-2 polypeptide.

The invention further provides a method of detecting a CLASP-2 polypeptide in a sample, comprising: (a) contacting the sample with a CLASP-2 polynucleotide or a polynucleotide that comprises a sequence of at least 12 nucleotides and is complementary to a contiguous sequence of the CLASP-2 polynucleotide and (b) determining whether a hybridization complex has been formed.

The invention further provides a method of detecting a CLASP-2 nucleotide in a sample, comprising: (a) using a polynucleotide that comprises a sequence of at least 12 nucleotides and is complementary to a contiguous sequence of a CLASP-2 polynucleotide in an amplification process; and (b) determining whether a specific amplification product has been formed.

The invention further provides pharmaceutical compositions comprising a CLASP-2 polynucleotide, a CLASP-2 polypeptide, or a CLASP-2 antibody and a pharmaceutically acceptable carrier.

In one aspect, the invention provides a method of inhibiting an immune response in a cell comprising: (a) interfering with the expression of a CLASP-2 gene in the cell; (b) interfering with the ability of a CLASP-2 protein to mediate cell-cell interaction (e.g., interfering with a heterotypic and/or homotypic interaction) between CLASP-2 and an extracellular protein; (c) interfering with the ability of a CLASP-2 protein to bind to another protein. In some such methods, the cell is a T cell or a B cell. Some such methods comprise contacting the cell with an effective amount of a polypeptide which comprises the amino acid sequence of SEQ ID NO: 2, 4, 6 or 10 or a fragment thereof.

In another aspect, the invention provides a method of inhibiting an immune response in a subject, comprising administering to the subject a therapeutically effective

amount of an antibody which specifically binds a polypeptide having the sequence of SEQ ID NO: 2, 4, 6 or 10.

In another aspect, the invention provides a method of preventing or treating a CLASP-2-mediated disease comprising administering to a subject in need thereof a therapeutically effective amount of a CLASP-2 pharmaceutical composition. In some such methods, the CLASP-2-mediated disease is an autoimmune disease.

The invention further provides a method of treating an autoimmune disease in a subject caused or exacerbated by increased activity of T<sub>H</sub>1 cells consisting of administering a therapeutically effective amount of a CLASP-2 pharmaceutical composition to the subject.

## BRIEF DESCRIPTION OF THE DRAWINGS

**Figure 1.** Nucleotide and predicted amino acid sequence of CLASP-2A cDNA. Notable protein motifs are indicated above the nucleotide sequence in bold. Potential initiator methionines are underscored. The notable, predicted protein motifs are: a cadherin cleavage site encoded by nucleotides 854-868, a cadherin ectodomain (EC) encoded by nucleotides 1253-1264, a transmembrane domain encoded by nucleotides 2861-2917, a coiled coil domain encoded by nucleotides 3579-3682, a second coiled coil domain encoded by nucleotides 3827-3937, and a PDZ binding motif (PBM) encoded by nucleotides 4046-4057.

**Figure 2. A.** Schematic of CLASP-2 splice variants. Splice variants are compared to Human (h) CLASP-2A. Numbers above hCLASP-2A line diagram indicate where splice variations comprising deletions and insertions relative to hCLASP-2A are found. Abbreviations: "KIAA" KIAA1058 sequence (Genbank Accession No. AB028981).

**B.** Nucleotide and predicted amino acid sequence of CLASP-2A cDNA. Notable protein motifs are indicated above the nucleotide sequence in bold. Exact position of insertions and deletions are indicated above the CLASP-2A sequence with arrows and "x", respectively.

The nucleotide sequence of insertions schematized in FIG. 2A are indicated above the arrow. The insertions and deletions are as follows (numeration refers to Human CLASP-2A nucleotide sequence): Nucleotides 1966-2034 are deleted in CLASP-2D. Nucleotides 2219-2224 are deleted in CLASP-2B. There is an insertion of 69 amino acids at nucleotide 2927 found in CLASP-2D. The nucleotide sequence for this insertion is:

AAGCAGTCCAGTGGGAGCCGCCCTTCTCCCCACAGCCATAGCGCCTGCCTGAG  
GAGGAGCCGGGGAG and encodes amino acids AVQWEPLLPHSHSACLRRSRG (one letter amino acid abbreviation). This amino acid sequence encodes a putative SH3 binding

domain. There is another deletion at between nucleotides 3011-3079 found in CLASP-2E. CLASPs 2B, 2C, 2D and 2E contain an insertion at nucleotide 3153 with the nucleotide sequence of:

TGAGAGGCTGGCCCATCTGTATGACACGCTGCACCGGGCCTACAGCAAAGTGAC  
 5 CGAGGTCATGCACTCGGGCCGCAGGCTTCTGGGGACCTACTTCCGGGGTAGCCTTC  
 TTCGGGCAGGCAGCGCAATACCAGTTTACAGACAGTGAAACAGATGTGGAGGGA  
 TT. The entire sequence is found in CLASP-2D and encodes amino acids  
 ERLAHL YDTLH RAYSKVTEVMHSGRRLLGTYFRVAFFGQAAQYQFTDSETDVEG

while the underline sequence is found in CLASPs 2B, 2C, and 2E and encodes amino acids  
 10 ERLAHL YDTLH RAYSKVTEVMHSGRRLLGTYFRVAFFGQG. This amino acid  
 sequence encodes a putative immunoreceptor tyrosine-based activation motif (ITAM). There  
 is a two nucleotide deletion in Human CLASP-2C found at nucleotides 3586 and 3587.  
 There is an insertion of 8 nucleotides found only in Human CLASP-2D with sequence:  
 CTGGGATG at nucleotide 3937. This insertion puts a stop codon into the CLASP-2D  
 15 nucleotide sequence.

**Figure 3. A.** Alignment of nucleotide sequences of the CLASP-2 isoforms. Sequences were aligned using ClustalW **B.** Alignment of amino acid sequence of the CLASP-2 isoforms. Sequences were aligned using ClustalW. One letter amino acid abbreviation is used.

**Figure 4.** Expression of CLASP-2 in human cell lines and human tissues as determined by Northern hybridization. A CLASP-2-specific DNA fragment was generated by PCR from a CLASP-2 cDNA clone (HC2-5'), using primers HC2AS2 and HC2S1. The fragment was labeled by incorporation of radioactive <sup>32</sup>P dCTP. **A.** Expression in human tissues. The labeled DNA fragment was used as a probe on a human Multiple Tissue Northern (Clontech MTN Blot, #7780-1). A single band is clearly detect migrating at approximately 7.5 kb in placenta, heart kidney and lung in the Multiple Tissue Northern. Slight expression is detected in liver, skeletal muscle and brain. **B.** Expression in hematopoietic cell lines. A Northern with RNA from multiple cells lines was hybridized with the same hCLASP-2 probe. A similarly migrating band is detected in Jurkat (T-cell derived),  
 20 9D10 (B-cell derived) and 293 (human kidney derived) cell lines. There are multiple weaker bands in the 9D10 lane indicating possible splice variants of hCLASP-2. Weak expression is  
 30 also detected in the mouse cell lines CH27 (B cell lymphoma) and 3A9 (T-cell hybridoma).



Since hybridization and washing were carried out at high stringency, this indicates that the human CLASP-2 probe may cross-react with mouse CLASP mRNA.

**Figure 5. A.** Amino acid sequence of human and rat CLASP proteins.

Sequences were aligned using ClustalW. One letter amino acid abbreviation used. Protein motifs are found within the labeled boxes. A “-” indicates gaps that are placed to acquire a best overall alignment. Other abbreviations: “HC2A” Human CLASP-2 sequence, “KIAA” KIAA1058 sequence (Genbank Accession No. AB028981), “rat” TRG gene (Genbank Accession No. X68101), “HC4” Human CLASP-4 sequence, “HC1” Human CLASP-1 sequence, “HC3” Human CLASP-3 sequence, “HC5” Human CLASP-5 sequence. **B.**

Alignment of DOCK motifs found within the human CLASPs and compared to canonical DOCK motifs. Consensus amino acids found within all DOCK motifs are also indicated.

**Figure 6. A.** Nucleotide and predicted amino acid sequence of CLASP-2A

cDNA. Notable protein motifs are indicated (see FIG. 1 legend for details). Additionally, boundaries between exons and introns are indicated by arrows. These boundaries were defined by sequencing Bacterial Artificial Chromosomes (BACs) containing genomic DNA corresponding to CLASP-2. BACs were sequenced using primers derived from exon sequences corresponding to the CLASP-2 cDNA. Each exon/intron boundary is noted (as “Ref” with an appropriate reference number) above the cDNA sequence. The references contain exact nucleotide location of introns. The names and nucleotide numbers of the primers that were used in sequence reactions are also indicated. All nucleotide numbers refer to CLASP-2A cDNA sequence. As shown in the reference, not all of the sequence from sequencing reactions produced sequence matching the cDNA. These nucleotide sequences that did not match the exon sequence for CLASP-2 were considered to be intron sequences.

**B.** Alignment of human and rat CLASP amino acid sequences by ClustalW. Notable protein motifs are indicated (see FIG. 1 Legend for additional details). Additionally, the exon/intron borders described in part A are indicated with vertical lines between appropriate amino acids. Reference numbers are indicated in the right margin and correspond to references in Fig 6A and B.

**Figure 7.** Southern hybridization analysis of CLASP-2. Genomic DNA from

HeLa cells or a BAC DNA clone was digested with EcoRI or HindIII (genomic DNA) or Pst I (BAC DNA) and electrophoresed and transferred to nylon membrane by standard methods. For a probe, a CLASP-2-specific DNA fragment was generated by PCR from a CLASP-2

cDNA clone (HC2-5'), using primers HC2AS2 and HC2S1. The fragment was labeled by incorporation of radioactive  $^{32}\text{P}$  dCTP. Probe HC2.1 is 800 bp long and it recognizes two fragments (~4.5 kb and 1.85 kb) on Eco RI digested genomic DNA. Three fragments are revealed by this probe when hybridized to digested DNA of BACs 4 and 6, with the two major ones identical in size to those detected on genomic DNA.

**Figure 8.** Expression of human CLASP-1 (hCLASP-1) CLASP-1 and CLASP-2 Glutathion-S-Transferase (GST) fusion proteins. Nucleotides encoding a portion of the hCLASP-2A intracellular domain (nucleotides 3230-4065) were subcloned into pGEX vectors (Pharmacia). Recombinant plasmids were transformed into *E. coli* (strain DH5 $\alpha$ ), and transformed strains were grown by standard conditions. While in log phase cells were either induced (I) with IPTG (0.1 mM final concentration) or left uninduced (U). After several additional hours of growth cells were harvested and soluble protein lysates generated by standard methods. Aliquots of the protein lysates were electrophoresed on SDS-PAGE along with molecular mass standards. The gel was stained with Coomassie Blue and shows that fusion proteins migrated with their predicted molecular masses of 59 and 57 kD for hCLASP-1 and hCLASP-2, respectively.

**Figure 9. A.** Binding of CLASP-2 C-terminal 20 amino acids to PDZ domains. 20  $\mu\text{M}$  biotinylated synthetic peptide corresponding to the C-terminal 20 amino acids of CLASP-2 was reacted with the indicated plate bound GST fusion proteins (none = no GST fusion protein coated onto plate). Error bars indicate standard deviation of duplicate measurements. **B.** Affinity of CLASP 2 – PDZ interactions. Varying concentrations of biotinylated CLASP-2 peptide were reacted with plate bound GST alone, GST-DLG1, GST-NeDLG, and GST-PSD95 fusion proteins. The binding to GST alone ( $< 0.1$  OD units) was subtracted from the binding to the fusion proteins and the remaining signal was divided by the signal observed upon addition of 30  $\mu\text{M}$  CLASP-2 peptide to each PDZ domain-containing protein (0.4 – 1.0 OD units) and plotted. The plotted data was fit to a saturation binding curve, yielding an apparent affinity of 7.5  $\mu\text{M}$  for NeDLG- CLASP-2 interaction, 21  $\mu\text{M}$  for DLG1- CLASP-2 interaction, and 45  $\mu\text{M}$  for PSD95-CLASP-2 interaction. Data are means of duplicate data points, with standard errors between duplicate data points  $< 10\%$ . **C.** Inhibition of CLASP-2 – PDZ binding. 5  $\mu\text{M}$  biotinylated synthetic peptide corresponding to the C-terminal 20 amino acids of CLASP-2 was reacted with the indicated, plate-bound PDZ domain-containing GST fusion proteins in the presence or absence of 100  $\mu\text{M}$  competitor



peptide. CLASP-2 Inhibitor refers to a synthetic peptide composed of the eight C-terminal amino acids of CLASP-2. KV1.3 Inhibitor refers to a synthetic peptide composed of the 19 C-terminal amino acids of KV1.3, a lymphocyte potassium channel. The amino acid sequence of the KV1.3 inhibitor is TTNNNPNSAVNIKKIFTDV. **D.** Inhibition of KV1.3 – PDZ binding. 5  $\mu$ M biotinylated synthetic peptide corresponding to the C-terminal 19 amino acids of KV1.3 was reacted with the indicated, plate-bound PDZ-domain containing GST fusion proteins in the presence or absence of 100  $\mu$ M CLASP-2 Inhibitor (see FIG. 9C legend).

**Figure 10.** Preliminary nucleotide sequences of CLASP-2 cDNAs.

**Figure 11.** A) Full length cDNA sequence and predicted amino acid translation of the human CLASP-2 gene. Predicted initiator methionine starts at nucleotide +1. Three independent 1st exons (indicated as 11A, 11B and 11C) splice into the second exon starting at nucleotide -101. The sequence appearing in FIG. 1 corresponds to nucleotides 1884 through 6690 of FIG 11A. **B)** Differences between the human CLASP-2 cDNA isoforms. In addition to the differential first exon usage indicated in **A**, sequencing multiple, independent cDNA products revealed nucleotide polymorphisms (allelic variations) between CLASP-2 cDNA isoforms. Additionally, differential exon usage through alternative splicing events was discovered. The use of the exon in **B** leads to a premature stop codon that can generate a soluble form of CLASP-2. **C.** Schematic of human CLASP-2 cDNAs. The top line represents nucleotide numbering found in FIG. 11A. Line (i) represents CLASP-2 cDNA shown in FIG. 1 above; line (ii) represents the full length CLASP-2 isoforms, where there are three CLASP-2 full length cDNA isoforms (A + Z, B + Z, and C + Z). Each of the isoforms uses a unique first exon (A, B, and C) (*see* FIG. 11A) that splices into the rest of the cDNA from exon 2 onwards represented by Z. The portion of the cDNA represented by Z itself has alternative splice and nucleotide polymorphisms that are shown in FIG. 2 above. Line (iii) represents the additional 5' sequence with a small region of overlap between nucleotides 1884 to 2109 in FIG. 11A and nucleotides 1-225 of FIG. 1.

**Figure 12.** Sequence of human CLASP-2 exons and intron boundaries. **A** Sequence of human CLASP-2 exons and intron borders. Stretches of noncontiguous genomic sequence from the Human Genome Project (GENBANK entry gi9988160) were aligned using the human CLASP-2 cDNA as a template and Sequencher sequence analysis software (Gene Codes Corp). 22 exons representing approximately the 5' 20% of the human CLASP-

2 cDNA sequence are presented in predicted 5' to 3' order. Exon sequences are underlined and are flanked by intron sequence. Nucleotide numbers in parentheses refer to the exon sequence within the uniquely-generated, contiguous gi9988160 sequence, which is located in

**B. B.** Ordered stretch of human genomic DNA at the CLASP-2 locus aligned from

5 noncontiguous, shotgun sequencing from the Human Genome Project using the human CLASP-2 sequence from FIG. 5A to determine genomic DNA fragment order and orientation.

**Figure 13.** Amino acid alignment and comparison between the human (h) CLASP family members. Amino acid sequences were aligned using ClustalW. The alignment is presented in order of their greatest pairwise similarity scores. Single letter amino acid abbreviations are used. Astericks indicate complete identity, while colons and periods indicate sequence similarity among CLASP family members. Dashes indicate gaps inserted in the amino acid sequence to facilitate alignment. Labelled boxes are domains with similarity to known protein motifs; unlabelled boxes represent regions of similarity between all CLASPs and may represent CLASP-specific domains.

**Figure 14.** Expression of CLASP-2 upon T-cell activation as assayed by Northern analysis. Jurkat cells were activated using PMA, Ionomycin, and  $\alpha$ CD28. RNA was prepared from cell culture aliquots at 0, 1, 2, 4, 8, 14 hours post activation and Northern analysis was performed (A). Hybridization signals obtained with a CLASP-2-specific probe were quantified using a phosphor imager system. Relative signal intensities (refers to total signal of the specific probe used) are shown in the bar diagram (B). The ethidium staining of the Northern gel (A) demonstrates even RNA loading.

## DETAILED DESCRIPTION

### 5.0 Definitions

25 Except when noted, the terms "patient" or "subject" are used interchangeably and refer to mammals such as human patients and non-human primates, as well as experimental animals such as rabbits, rats, and mice, and other animals.

The term "biological sample" as used herein is a sample of biological tissue, fluid, or cells that contains hCLASP-2 or nucleic acid encoding hCLASP-2 protein. Such samples include, but are not limited to, tissue isolated from humans. Biological samples may also include sections of tissues such as frozen sections taken for histologic purposes. A biological sample is typically obtained from a eukaryotic organism, preferably eukaryotes

such as fungi, plants, insects, protozoa, birds, fish, reptiles, and preferably a mammal such as rat, mice, cow, dog, guinea pig, or rabbit, and most preferably a primate such as chimpanzees or humans.

The term “treating” includes the administration of the compounds or agents of the present invention to prevent or delay the onset of the symptoms, complications, or biochemical indicia of a disease, alleviating the symptoms or arresting or inhibiting further development of the disease, condition, or disorder (*e.g.*, autoimmune disease). Treatment may be prophylactic (to prevent or delay the onset of the disease, or to prevent the manifestation of clinical or subclinical symptoms thereof) or therapeutic suppression or alleviation of symptoms after the manifestation of the disease.

The term “lymphocyte” as used herein has the normal meaning in the art, and refers to any of the mononuclear, nonphagocytic leukocytes, found in the blood, lymph, and lymphoid tissues, *i.e.*, B and T lymphocytes.

The terms “isolated,” or “purified,” refer to material that is substantially free from components that normally accompany it as found in its native state (*e.g.*, recombinantly produced or purified away from other cell components with which it is naturally associated). Purity and homogeneity are typically determined using analytical chemistry techniques such as polyacrylamide gel electrophoresis or high performance liquid chromatography. The term “purified” denotes that a nucleic acid or protein gives rise to essentially one band in an electrophoretic gel. Particularly, it means that the nucleic acid or protein is at least 85% pure, more preferably at least 95% pure, and most preferably at least 99% pure.

The terms “nucleic acid” and “polynucleotide” are used interchangeably and refer to refers to DNA, RNA and nucleic acid polymers containing known nucleotide analogs or modified backbone residues or linkages, which are synthetic, naturally occurring, and non-naturally occurring, which have similar binding properties as the reference nucleic acid, and which are metabolized in a manner similar to the reference nucleotides. Examples of such analogs include, without limitation, phosphorothioates, phosphoramidates, methyl phosphonates, chiral-methyl phosphonates, 2-O-methyl ribonucleotides, peptide-nucleic acids (PNAs).

The terms “polypeptide,” “peptide” and “protein” are used interchangeably herein to refer to a polymer of amino acid residues. The amino acids may be natural amino acids, or include an artificial chemical mimetic of a corresponding naturally occurring amino acid.

As used herein a “nucleic acid probe” is defined as a nucleic acid capable of specifically binding to a target nucleic acid of complementary sequence (*e.g.*, through complementary base pairing). As used herein, a probe may include natural (*i.e.*, A, G, C, or T) or modified bases (7-deazaguanosine, inosine, and the like). In addition, the bases in a probe may be joined by a linkage other than a phosphodiester bond, so long as it does not interfere with hybridization (*e.g.*, probes may be peptide nucleic acids). The probes can be directly labeled as with isotopes, chromophores, lumiphores, chromogens, or indirectly labeled such as with biotin to which a streptavidin complex may later bind.

The term “recombinant” when used with reference, *e.g.*, to a cell, or nucleic acid, protein, or vector, indicates that the cell, nucleic acid, protein or vector, has been modified by the introduction of a heterologous nucleic acid or protein or the alteration of a native nucleic acid or protein, or, in the case of cells, to progeny of a cell so modified. Thus, for example, recombinant cells express genes that are not found within the native (non-recombinant) form of the cell or express native genes that are otherwise abnormally expressed, under expressed or not expressed at all.

The term “heterologous” when used with reference to portions of a nucleic acid indicates that the nucleic acid comprises two or more subsequences that are not found in the same relationship to each other in nature. For instance, the nucleic acid is typically recombinantly produced, having two or more sequences from unrelated genes arranged to make a new functional nucleic acid, *e.g.*, a promoter from one source and a coding region from another source. Similarly, a heterologous protein indicates that the protein comprises two or more subsequences that are not found in the same relationship to each other in nature (*e.g.*, a fusion protein).

The term “sequence identity” refers to a measure of similarity between amino acid or nucleotide sequences, and can be measured using methods known in the art, such as those described below:

The terms “identical” or percent “identity,” in the context of two or more nucleic acids or polypeptide sequences, refer to two or more sequences or subsequences that are the same or have a specified percentage of amino acid residues or nucleotides that are the same (*i.e.*, 60% identity, preferably 65%, 70%, 75%, 80%, 85%, 90%, or 95% identity over a specified region (see, *e.g.*, SEQ ID NO: 1 ), when compared and aligned for maximum correspondence over a comparison window, or designated region as measured using one of the following sequence comparison algorithms or by manual alignment and visual inspection.

The phrase “substantially identical,” in the context of two nucleic acids or polypeptides, refers to two or more sequences or subsequences that have at least of at least 60%, often at least 70%, preferably at least 80%, most preferably at least 90% or at least 95% nucleotide or amino acid residue identity, when compared and aligned for maximum correspondence, as measured using one of the following sequence comparison algorithms or by visual inspection. Preferably, the substantial identity exists over a region of the sequences that is at least about 50 bases or residues in length, more preferably over a region of at least about 100 bases or residues, and most preferably the sequences are substantially identical over at least about 150 bases or residues. In a most preferred embodiment, the sequences are substantially identical over the entire length of the coding regions.

The phrase “sequence similarity” in the context of two nucleic acids or polypeptides, refers to two or more sequences that are identical or in the case of amino acids, have homologous amino acid substitutions at either 50%, often at least 60%, often at least 70%, preferably at least 80%, most preferably at least 90% or at least 95% of the indicated positions.

For sequence comparison, typically one sequence acts as a reference sequence, to which test sequences are compared. When using a sequence comparison algorithm, test and reference sequences are entered into a computer, subsequence coordinates are designated, if necessary, and sequence algorithm program parameters are designated. Default program parameters can be used, or alternative parameters can be designated. The sequence comparison algorithm then calculates the percent sequence identities for the test sequences relative to the reference sequence, based on the program parameters. For sequence comparison of nucleic acids and proteins to CLASP-2 nucleic acids and proteins, the BLAST and BLAST 2.0 algorithms and the default parameters discussed below are used.

A “comparison window”, as used herein, includes reference to a segment of any one of the number of contiguous positions selected from the group consisting of from 20 to 600, usually about 50 to about 200, more usually about 100 to about 150 in which a sequence may be compared to a reference sequence of the same number of contiguous positions after the two sequences are optimally aligned. Methods of alignment of sequences for comparison are well-known in the art. Optimal alignment of sequences for comparison can be conducted, *e.g.*, by the local homology algorithm of Smith & Waterman, 1981, *Adv. Appl. Math.* 2: 482), by the homology alignment algorithm of Needleman & Wunsch, 1970, *J. Mol. Biol.* 48: 443, by the search for similarity method of Pearson & Lipman, 1988, *Proc. Natl. Acad. Sci. U.S.A.* 85: 2444, by computerized implementations of these algorithms



(FASTDB (Intelligenetics), BLAST (National Center for Biomedical Information), GAP, BESTFIT, FASTA, and TFASTA in the Wisconsin Genetics Software Package, Genetics Computer Group, 575 Science Dr., Madison, WI), or by manual alignment and visual inspection (see, *e.g.*, Ausubel *et al.*, 1987 (1999 Suppl.), Current Protocols in Molecular Biology, Greene Publishing Associates and Wiley Interscience, N.Y.)

A preferred example of an algorithm that is suitable for determining percent sequence identity and sequence similarity is the FASTA algorithm, which is described in Pearson, W.R. & Lipman, D.J., 1988, Proc. Natl. Acad. Sci. U.S.A. 85: 2444. See also W. R. Pearson, 1996, Methods Enzymol. 266: 227-258. Preferred parameters used in a FASTA alignment of DNA sequences to calculate percent identity are optimized, BL50 Matrix 15: -5, k-tuple= 2; joining penalty= 40, optimization= 28; gap penalty -12, gap length penalty =-2; and width= 16.

Another preferred example of algorithm that is suitable for determining percent sequence identity and sequence similarity are the BLAST and BLAST 2.0 algorithms, which are described in Altschul *et al.*, 1977, Nuc. Acids Res. 25: 3389-3402 and Altschul *et al.*, 1990, J. Mol. Biol. 215: 403-410, respectively. BLAST and BLAST 2.0 are used, with the parameters described herein, to determine percent sequence identity for the nucleic acids and proteins of the invention. Software for performing BLAST analyses is publicly available through the National Center for Biotechnology Information (<http://www.ncbi.nlm.nih.gov/>). This algorithm involves first identifying high scoring sequence pairs (HSPs) by identifying short words of length W in the query sequence, which either match or satisfy some positive-valued threshold score T when aligned with a word of the same length in a database sequence. T is referred to as the neighborhood word score threshold (Altschul *et al.*, *supra*). These initial neighborhood word hits act as seeds for initiating searches to find longer HSPs containing them. The word hits are extended in both directions along each sequence for as far as the cumulative alignment score can be increased. Cumulative scores are calculated using, for nucleotide sequences, the parameters M (reward score for a pair of matching residues; always > 0) and N (penalty score for mismatching residues; always < 0). For amino acid sequences, a scoring matrix is used to calculate the cumulative score. Extension of the word hits in each direction are halted when: the cumulative alignment score falls off by the quantity X from its maximum achieved value; the cumulative score goes to zero or below, due to the accumulation of one or more negative-scoring residue alignments; or the end of either sequence is reached. The BLAST algorithm parameters W, T, and X determine the sensitivity and speed of the alignment. The BLASTN program (for nucleotide sequences)

uses as defaults a wordlength (W) of 11, an expectation (E) of 10, M=5, N=-4 and a comparison of both strands. For amino acid sequences, the BLASTP program uses as defaults a wordlength of 3, and expectation (E) of 10, and the BLOSUM62 scoring matrix (see Henikoff & Henikoff, 1989, Proc. Natl. Acad. Sci. U.S.A. 89: 10915) alignments (B) of 50, expectation (E) of 10, M=5, N=-4, and a comparison of both strands.

The BLAST algorithm also performs a statistical analysis of the similarity between two sequences (see, e.g., Karlin & Altschul, 1993, Proc. Natl. Acad. Sci. U.S.A. 90: 5873-5787). One measure of similarity provided by the BLAST algorithm is the smallest sum probability (P(N)), which provides an indication of the probability by which a match between two nucleotide or amino acid sequences would occur by chance. For example, a nucleic acid is considered similar to a reference sequence if the smallest sum probability in a comparison of the test nucleic acid to the reference nucleic acid is less than about 0.2, more preferably less than about 0.01, and most preferably less than about 0.001.

Another example of a useful algorithm is PILEUP. PILEUP creates a multiple sequence alignment from a group of related sequences using progressive, pairwise alignments to show relationship and percent sequence identity. It also plots a tree or dendrogram showing the clustering relationships used to create the alignment. PILEUP uses a simplification of the progressive alignment method of Feng & Doolittle, 1987, J. Mol. Evol. 35: 351-360. The method used is similar to the method described by Higgins & Sharp, 1989, CABIOS 5: 151-153. The program can align up to 300 sequences, each of a maximum length of 5,000 nucleotides or amino acids. The multiple alignment procedure begins with the pairwise alignment of the two most similar sequences, producing a cluster of two aligned sequences. This cluster is then aligned to the next most related sequence or cluster of aligned sequences. Two clusters of sequences are aligned by a simple extension of the pairwise alignment of two individual sequences. The final alignment is achieved by a series of progressive, pairwise alignments. The program is run by designating specific sequences and their amino acid or nucleotide coordinates for regions of sequence comparison and by designating the program parameters. Using PILEUP, a reference sequence is compared to other test sequences to determine the percent sequence identity relationship using the following parameters: default gap weight (3.00), default gap length weight (0.10), and weighted end gaps. PILEUP can be obtained from the GCG sequence analysis software package, e.g., version 7.0 (Devereaux *et al.*, 1984, Nuc. Acids Res. 12: 387-395).

Another preferred example of an algorithm that is suitable for multiple DNA and amino acid sequence alignments is the CLUSTALW program (Thompson, J. D. *et al.*,



1994, Nucl. Acids. Res. 22: 4673-4680). ClustalW performs multiple pairwise comparisons between groups of sequences and assembles them into a multiple alignment based on homology. Gap open and Gap extension penalties were 10 and 0.05 respectively. For amino acid alignments, the BLOSUM algorithm can be used as a protein weight matrix (Henikoff and Henikoff, 1992, Proc. Natl. Acad. Sci. U.S.A. 89: 10915-10919).

A "label" is a composition detectable by spectroscopic, photochemical, biochemical, immunochemical, or chemical means. For example, useful labels include  $^{32}\text{P}$ , fluorescent dyes, electron-dense reagents, enzymes (*e.g.*, as commonly used in an ELISA), biotin, digoxigenin, or haptens and proteins for which antisera or monoclonal antibodies are available (*e.g.*, the polypeptide of SEQ ID NO: 1 can be made detectable, *e.g.*, by incorporating a radiolabel into the peptide, and used to detect antibodies specifically reactive with the peptide).

The term "sorting" in the context of cells as used herein to refers to both physical sorting of the cells, as can be accomplished using, *e.g.*, a fluorescence activated cell sorter, as well as to analysis of cells based on expression of cell surface markers, *e.g.*, FACS analysis in the absence of sorting.

The phrase "selectively (or specifically) hybridizes to" refers to the binding, duplexing, or hybridizing of a molecule only to a particular nucleotide sequence under stringent hybridization conditions when that sequence is present in a complex mixture (*e.g.*, total cellular or library DNA or RNA).

The phrase "specifically (or selectively) binds" to an antibody refers to a binding reaction that is determinative of the presence of the protein in a heterogeneous population of proteins and other biologics. Thus, under designated immunoassay conditions, the specified antibodies bind to a particular protein at least two times the background and do not substantially bind in a significant amount to other proteins present in the sample.

The phrase "specifically bind(s)" or "bind(s) specifically" when referring to a peptide refers to a peptide molecule which has intermediate or high binding affinity, exclusively or predominately, to a target molecule. The phrases "specifically binds to" refers to a binding reaction which is determinative of the presence of a target protein in the presence of a heterogeneous population of proteins and other biologics. Thus, under designated assay conditions, the specified binding moieties bind preferentially to a particular target protein and do not bind in a significant amount to other components present in a test sample. Specific binding to a target protein under such conditions may require a binding moiety that is selected for its specificity for a particular target antigen. A variety of assay formats may be

used to select ligands that are specifically reactive with a particular protein. For example, solid-phase ELISA immunoassays, immunoprecipitation, Biacore and Western blot are used to identify peptides that specifically react with PDZ domain-containing proteins. Typically a specific or selective reaction will be at least twice background signal or noise and more typically more than 10 times background. Specific binding between a monovalent peptide and a PDZ-containing protein means a binding affinity of at least  $10^4 \text{ M}^{-1}$ , and preferably  $10^5$  or  $10^6 \text{ M}^{-1}$ .

The phrase “homotypic interaction” refers to the binding of a given protein to another molecule of the same protein (*e.g.*, the binding of hCLASP-2 to hCLASP-2). The phrase “heterotypic interaction” refers to the binding of a given protein to a different protein or other molecule (*e.g.*, the binding of hCLASP-2 to a PDZ domain-containing protein or the binding of a transcription factor to DNA).

The phrase “immune cell response” refers to the response of immune system cells to external or internal stimuli (*e.g.*, antigen, cytokines, chemokines, and other cells) producing biochemical changes in the immune cells that result in immune cell migration, killing of target cells, phagocytosis, production of antibodies, other soluble effectors of the immune response, and the like.

The terms “B lymphocyte response” and “B lymphocyte activity” are used interchangeably to refer to the component of immune response carried out by B lymphocytes (*i.e.* the proliferation and maturation of B lymphocytes, the binding of antigen to cell surface immunoglobulin, the internalization of antigen and presentation of that antigen via MHC molecules to T lymphocytes, and the synthesis and secretion of antibodies).

The terms “T lymphocyte response” and “T lymphocyte activity” are used here interchangeably to refer to the component of immune response dependent on T lymphocytes (*i.e.*, the proliferation and/or differentiation of T lymphocytes into helper, cytotoxic killer, or suppressor T lymphocytes, the provision of signals by helper T lymphocytes to B lymphocytes that cause or prevent antibody production, the killing of specific target cells by cytotoxic T lymphocytes, and the release of soluble factors such as cytokines that modulate the function of other immune cells).

The term “immune response” refers to the concerted action of lymphocytes, antigen presenting cells, phagocytic cells, granulocytes, and soluble macromolecules produced by the above cells or the liver (including antibodies, cytokines, and complement) that results in selective damage to, destruction of, or elimination from the human body of

invading pathogens, cells or tissues infected with pathogens, cancerous cells, or, in cases of autoimmunity or pathological inflammation, normal human cells or tissues.

Components of an immune response may be detected in vitro by various methods that are well known to those of ordinary skill in the art. For example, (1) cytotoxic T lymphocytes can be incubated with radioactively labeled target cells and the lysis of these target cells detected by the release of radioactivity, (2) helper T lymphocytes can be incubated with antigens and antigen presenting cells and the synthesis and secretion of cytokines measured by standard methods (Windhagen A; *et al.*, 1995, Immunity 2(4): 373-80), (3) antigen presenting cells can be incubated with whole protein antigen and the presentation of that antigen on MHC detected by either T lymphocyte activation assays or biophysical methods (Harding *et al.*, 1989, Proc. Natl. Acad. Sci., 86: 4230-4), (4) mast cells can be incubated with reagents that cross-link their Fc-epsilon receptors and histamine release measured by enzyme immunoassay (Siraganian, *et al.*, 1983, TIPS 4: 432-437).

Similarly, products of an immune response in either a model organism (*e.g.*, mouse) or a human patient can also be detected by various methods that are well known to those of ordinary skill in the art. For example, (1) the production of antibodies in response to vaccination can be readily detected by standard methods currently used in clinical laboratories, *e.g.*, an ELISA; (2) the migration of immune cells to sites of inflammation can be detected by scratching the surface of skin and placing a sterile container to capture the migrating cells over scratch site (Peters *et al.*, 1988, Blood 72: 1310-5); (3) the proliferation of peripheral blood mononuclear cells in response to mitogens or mixed lymphocyte reaction can be measured using <sup>3</sup>H-thymidine; (4) the phagocytic capacity of granulocytes, macrophages, and other phagocytes in PBMCs can be measured by placing PMBCs in wells together with labeled particles (Peters *et al.*, 1988); and (5) the differentiation of immune system cells can be measured by labeling PBMCs with antibodies to CD molecules such as CD4 and CD8 and measuring the fraction of the PBMCs expressing these markers.

As used herein, the phrase "signal transduction pathway" or "signal transduction event" refers to at least one biochemical reaction, but more commonly a series of biochemical reactions, which result from interaction of a cell with a stimulatory compound or agent. Thus, the interaction of a stimulatory compound with a cell generates a "signal" that is transmitted through the signal transduction pathway, ultimately resulting in a cellular response, *e.g.*, an immune response described above.

A signal transduction pathway refers to the biochemical relationship between a variety of signal transduction molecules that play a role in the transmission of a signal from

one portion of a cell to another portion of a cell. Signal transduction molecules of the present invention include, for example, extracellular and intracellular domains of CLASP-2. As used herein, the phrase "cell surface receptor" includes molecules and complexes of molecules capable of receiving a signal and the transmission of such a signal across the plasma membrane of a cell. An example of a "cell surface receptor" of the present invention is the T cell receptor (TCR). As used herein, the phrase "intracellular signal transduction molecule" includes those molecules or complexes of molecules involved in transmitting a signal from the plasma membrane of a cell through the cytoplasm of the cell, and in some instances, into the cell's nucleus. In the present invention, CLASP-2 can be referred to as an "intracellular signal transduction molecule", but can also be referred to as a "signal transduction molecule".

A signal transduction pathway in a cell can be initiated by interaction of a cell with a stimulator that is inside or outside of the cell. If an exterior (*i.e.*, outside of the cell) stimulator (*e.g.*, an MHC-antigen complex on an antigen presenting cell) interacts with a cell surface receptor (*e.g.*, a T cell receptor), a signal transduction pathway can transmit a signal across the cell's membrane, through the cytoplasm of the cell, and in some instances into the nucleus. If an interior (*e.g.*, inside the cell) stimulator interacts with an intracellular signal transduction molecule, a signal transduction pathway can result in transmission of a signal through the cell's cytoplasm, and in some instances into the cell's nucleus.

Signal transduction can occur through, *e.g.*, the phosphorylation of a molecule; non-covalent allosteric interactions; complexing of molecules; the conformational change of a molecule; calcium release; inositol phosphate production; proteolytic cleavage; cyclic nucleotide production and diacylglyceride production. Typically, signal transduction occurs through phosphorylating a signal transduction molecule. According to the present invention, a CLASP-2 signal transduction pathway refers generally to a pathway in which CLASP-2 protein regulates a pathway that includes engaged-receptors, PKC-substrates, G proteins, and other molecules.

### **5.1. Introduction**

The present invention relates to a novel transmembrane protein, CLASP-2, a new member of the CLASP family that contains an endodomain that displays the appropriate properties to organize the cytoskeleton and signal transduction apparatus of the immune gateway.

CLASP-2 functions in cells of the immune system, *e.g.*, T cells and B cells, as well as non-immune cells. The CLASP-2 protein functions in a variety of cellular processes,

particularly related to immune function, regulation of T cell and B cell interactions, T cell activation, and in the organization, establishment and maintenance of the “immunological synapse” (see Dustin *et al.*, 1999, Science 283: 680-682; Paul *et al.*, 1994, Cell 76: 241-251; Dustin *et al.*, 1996, J. Immunol. 157: 2014; Dustin *et al.*, 1998, Cell 94: 667), including  
5 signal transduction, cytoskeletal interactions, and membrane organization.

Without intending to be bound by a particular mechanism or limited in any way, the CLASP-2 protein is believed to be a component of the lymphocyte organelle called the “immune gateway” that creates a docking site or portal for cell-cell contact during antigen-presentation. It is believed the cytoplasmic domains of CLASP-2 proteins organize it  
10 into a patch at the leading edge of T cells. The carboxy-terminus encoded sequences mediate interaction with PDZ domain proteins and with cytoskeletal proteins (*e.g.*, spectrin or ankyrin) to connect CLASP-2 to the microtubule network and hold the receptors at a polarized configuration just above the microtubule-organizing center (“MTOC”). Thus, when T cells engages a B cell acting as an APC, the CLASP-2 molecules engage one another  
15 to dock the two cells and organize the immune synapse.

Modulating the expression of the CLASP-2 protein, and interference with, or enhancement of, CLASP-2 protein interactions with other proteins has a number of beneficial physiological effects, *e.g.*, altered signaling in response to antigen, altered T and B cell response to antigen, and modulation of T cell activation. In one aspect, the CLASP-2  
20 extracellular domain is targeted (*e.g.*, using anti-CLASP-2 antibody, soluble CLASP-2 fragments, and the like) to regulate T cell activation (and thus regulate immune responses). Disorders that can be treated by disrupting CLASP-2 function, include without limitation, multiple sclerosis, juvenile diabetes, rheumatoid arthritis, pemphigus, pemphigoid, epidermolysis bullosa acquista, lupus, endometriosis, toxemia or pregnancy induced  
25 hypertension, pruritic urticarial papules and plaques of pregnancy (PUPPP), herpes gestationis, impetigo herpetiformis, pruritus gravidarum, placenta-related disorders, and Rh incompatibility.

In another aspect, the present invention provides methods and reagents for detection of CLASP-2 expression and CLASP-2-expressing cells. Abnormal expression  
30 patterns or expression levels are diagnostic for immune and other disorders. For example, diseases characterized by overproduction or depletion of lymphocytes in blood or other organs may be detected or monitored by monitoring the level of CLASP-2 polypeptide or mRNA in a biological sample (*e.g.*, peripheral blood), *e.g.*, the number or percentage of CLASP-2 expressing cells. Diseases characterized by overproduction of T cells include, *e.g.*,



leukemia (both ALL and CLL), lymphoma (including non-Hodgkins lymphoma, Burkitt's lymphoma, mycosis fungoides, and sezary syndrome), EBV, CMV, toxoplasmosis, syphilis, typhoid, brucellosis, tuberculosis, influenza, hepatitis, serum sickness, and thyrotoxicosis. Diseases associated with the depletion of T cells include, *e.g.*, HIV and myelodysplasia.

5 Diseases associated with the overproduction of B cells include, *e.g.*, leukemia (both ALL and CLL), non-Hodgkins lymphoma, Burkitt's lymphoma, myeloma, EBV, CMV, toxoplasmosis, syphilis, typhoid, brucellosis, tuberculosis, influenza, hepatitis, serum sickness, and thyrotoxicosis. Diseases associated with the depletion of B cells include, *e.g.*, myelodysplasia.

## 10 **5.2. CLASP-2 cDNA and Polypeptide Structure**

The CLASP-2 protein is type I transmembrane glycoprotein, characterized by multiple forms produced by alternative exon usage (*i.e.*, production of splice variants). In one naturally occurring form, CLASP-2 has the structure shown in FIG. 1. However, as discussed in detail *infra*, the CLASP-2 gene encodes a variety of gene product due to  
15 alternative splicing of mRNA. FIG. 2 shows the nucleotide sequence and conceptual translation of human CLASP-2 polypeptides:

hCLASP-2A cDNA (SEQ ID NO: 1) and hCLASP-2A polypeptide (SEQ ID NO: 2).

hCLASP-2B cDNA (SEQ ID NO: 3) and hCLASP-2B polypeptide (SEQ ID NO: 4).

hCLASP-2C cDNA (SEQ ID NO: 5) and hCLASP-2C polypeptide (SEQ ID NO: 6).

hCLASP-2D cDNA (SEQ ID NO: 7) and hCLASP-2D polypeptide (SEQ ID NO: 8).

hCLASP-2E cDNA (SEQ ID NO: 9) and hCLASP-2E polypeptide (SEQ ID NO: 10).

Unless specifically referred to, the phrase "human CLASP-2 (hCLASP-2)" is used herein refers to hCLASP-2A, hCLASP-2B, hCLASP-2C and hCLASP-2E. "hCLASP-2D" cDNA is also known as KIAA1058, which was described by Kikuno *et al.*, 1999, *DNA Res.* 6, 197-205 as a cDNA from brain encoding a protein of unknown function.

CLASP-2 polypeptides typically include an approximately 120 residue leader sequence, followed by a cadherin proteolytic cleavage signal RXXR, an extracellular domain, a transmembrane domain, and an intracellular domain. The present invention provides a polynucleotide having the sequence of SEQ. ID. NO: 1, or a fragment thereof, and a  
35 polypeptide having the sequence of SEQ. ID NO: 2, or a fragment thereof. In addition, the

invention provides polynucleotides comprising hCLASP-2 genomic sequences, CLASP-2 homologs from other species, naturally occurring alleles of hCLASP-2, and hCLASP-2 variants as described herein, and methods for using CLASP-2 polynucleotide, polypeptides, antibodies and other reagents.

### 5 5.2.1. CLASP-2 Polypeptide Domains

As is shown in FIG. 1, one naturally occurring CLASP-2 cDNA encodes a polypeptide characterized by several structural and functional domains and defined sequence motifs. To provide guidance to the practitioner, the structural features are described *infra*. However, it will be understood that the present invention is not limited to polypeptides that include all, or any particular one of these domains or motifs. For example, a CLASP-2 fusion protein of the invention contains only the extracellular domain of CLASP-2. Similarly, the CLASP-2A polypeptide of SEQ ID NO: 2 does not have the ITAM motifs (discussed *infra*) found in the CLASP-2B and 2C polypeptides.

It will be appreciated that the structurally (and functionally) different domains of CLASP-2 polypeptides (and the corresponding region of the mRNA) are of interest, in part, because they may be separately targeted or modified (*e.g.*, deleted or mutated) to affect the activity or expression of a CLASP-2 gene product (in order to, for example, modulate an immune response). For example, the extracellular domain of a CLASP-2 protein can be targeted (*e.g.*, using an anti-CLASP monoclonal antibody to (a) block the interaction of a CLASP-2-expressing cell (*e.g.*, a T cell) and a second cell (*e.g.*, a B cell) displaying a protein that is bound by CLASP-2 (*i.e.*, a CLASP-2 ligand). Similarly, an intracellular domain (*e.g.*, ITAM or DOCK, see *infra*) can be targeted to interfere with signal transduction without interfering with extracellular ligand binding.

Generally, inhibiting CLASP-2 expression or CLASP-2 polypeptide function will result in modulation of immune function including, for example, changing the threshold for T cell activation by affecting formation of the immune synapse. Modulation of immune function can be screened and quantitated by a number of assays known in the art and described herein (see also §5.14).

#### 5.2.1.1. Signal Peptide

The human CLASP-2 sequence presented in FIG. 1 encodes two potential start sites for translation. The first predicted methionine appears at nucleotide 278 (ATG). The second methionine appears at nucleotide 476. Both have an acceptable consensus sequence



for a translational start (A/GxxATGG; Kozak, M., 1996, Mamm. Genome 7(8): 563-74 ). A polypeptide beginning at the second methionine is also predicted to encode a signal peptide capable of localizing the protein to the secretory pathway by SignalP, a signal sequence prediction program (Nielsen, H. *et al.*, 1997, Protein Eng. 10(1): 1-6). Polypeptides beginning at the first methionine are not predicted to contain a signal sequence; however, the consensus for signal prediction is only 80-90% accurate for known signal sequences. A third possibility for a translational start is that the cDNA listed in FIG. 1 is incomplete and another methionine is encoded in frame and upstream of the sequence shown in FIG.1 .

#### **5.2.1.2. Extracellular Domain**

The CLASP-2 extracellular domain is characterized by one cadherin EC-like motif (Pigott, R. and Power, C., 1993, The Adhesion Molecule Factbook. Academic Press, pg. 6; Jackson, R. M. and Russell, R. B., 2000, J. Mol. Biol. 296: 325-34). Several highly conserved cysteines are found in the extracellular domain, as well as various glycosylation signals. Through its extracellular domains, CLASP-2 may interact with ligands in a homotypic and/or heterotypic manner to establish the immunological synapse in conjunction with molecules such as TCR, MHC class I, MHC class II, CD3 complex and accessory molecules such as CD4, CD3, ICAM-1, LFA-1, and others. Many cadherins contain a pro-domain of approximately 50 to 150 amino acids that is removed before localization to the plasma membrane. This cleavage is presumed to be carried out by Furin (Posthaus, H. *et al.*, 1998, FEBS Let 438: 306-10) at a consensus sequence of RKQR. Furin is a protease that is at least partially responsible for the maturation of certain cadherins. CLASP-2 has the sequence RNQR at nucleotides 945 through 957. By homology, this region is around 120 amino acids into the predicted protein start site for hCLASP-2A. This region may be a pro-domain and cleavage may be required for CLASP-2 function, or aspects of CLASP-2 function.

Antibodies raised against the extracellular domain can be added to cells expressing CLASP-2. These antibodies can either block the interaction of CLASP-2 with potential ligands or stabilize these interactions. Any immunoassay known in the art, *e.g.*, listed and described herein, may be used to assess the modulation of immune function brought about by this approach.

Similarly, portions of the extracellular domain of CLASP-2 can be expressed as soluble protein. This soluble protein can then be added to cells expressing CLASP-2. These proteins may interact with potential ligands to competitively inhibit their binding to endogenous CLASP-2. This could modulate CLASP-2 function via the immunoassays

described herein. Recombinant proteins could interfere in a positive or negative fashion with CLASP-2 interactions.

#### **5.2.1.3. Transmembrane Domain**

CLASP-2 predicted amino acid sequence was analyzed using the PHDhtm analysis software for prediction of transmembrane helices (Rost, B., *et al.*, 1996, *Prot. Science* 7: 1704-1718). Using the PPHDhtm analysis software, it was determined that the transmembrane domain is located from nucleotides 2861-2917 (see FIG. 1), as well as three other potential transmembrane domains located near the amino terminal end.

#### **5.2.1.4. Intracellular Domains**

The CLASP-2 intracellular domains contain motifs corresponding to several types of protein domains. Depending on the specific CLASP-2 (*i.e.*, specific family member or splice variant) all or only some of the domains can be present. Listed from amino terminus to carboxy terminus, the domains include: (1) ITAM (Chan *et al.* 1994, *Annual Review of Immunology* 12: 555-592), (2) a newly discovered DOCK/CLASP-2 motif, (3) a coiled-coil motif, and (4) a C-terminal PDZ binding motif (PBM) (also referred to as PDZ ligand or "PL").

#### **5.2.1.5. ITAM**

Immunoreceptor Tyrosine-based Activation Motifs (ITAM motifs; also known as ARAM, or antigen recognition activation motifs) are motifs contained within antigen receptors for T and B cells, and Fc receptors on other leukocytes, and are necessary for proper activation and signal transduction in these cells. They are characterized by the consensus sequence YXXL/I - X7/8- YXXL/I (Grucza *et al.*, 1999, *Biochemistry* 38: 5024-5033), usually separated by 6-8 amino acids (Watson *et al.*, 1998, *Immunol. Today* 19: 260-264; Isakov, J. *Leukoc. Biol.* 61: 6-16). ITAM is used as an intracellular regulatory motif through its ability to be tyrosine phosphorylated by src-family tyrosine kinases such as Lyn that are involved in leukocyte signal transduction. Once phosphorylated, the ITAM acts as a high affinity binding site for SH2 containing proteins. Signal transduction components including ZAP-70, Syk, Lyn, Shc, PI3 kinase, and Grb2 contain SH2 domains and have been shown to bind ITAMs (Clements *et al.*, 1999, *Annu. Rev. Immunol.* 17: 89-108). This places ITAM-containing molecules in a central role of intracellular signal regulation in leukocytes. ITAM motifs in leukocyte signaling can facilitate signal transduction (*e.g.*,

tyrosine kinase signaling) by acting as temporal scaffolds where other transduction components could bind and be properly positioned to mediate transduction. ITAM motifs often appear in multiples in a protein, however, it is known that one set of YXXL/I alone can transduce signals of the PTK pathway, though weakly.

CLASP-2 proteins typically have ITAM YXXL/I motifs (where X is any amino acid) separated by 3 or 13 amino acids. In various embodiments the CLASP-2 polypeptide of the invention is characterized by one or more of the motifs shown in Table 1.

**Table 1**  
**CLASP-2 ITAM Motifs**

Motif No.	Sequence Motif
1	YXX(I/L)-X <sub>3</sub> -YXX(I/L)
2	YXX(I/L)-X <sub>13</sub> -YXX(I/L)
3	YXX(I/L)-X <sub>3</sub> -YXX(I/L)-X <sub>13</sub> -YXX(I/L)

The presence of multiple ITAM motifs in CLASPs proteins indicates that they may be engaged by multiple signal transduction components (*e.g.*, ZAP-70/Syk, Shc, PI3 kinase, and Grb2). In general, the ITAM motif in CLASP proteins match identically to the canonical ITAM motif with some motifs containing a conservative amino acid change (*i.e.* valine instead of isoleucine or leucine). As previously described for other ITAMs, the ITAMs within CLASPs can bind SH2-containing proteins including ZAP-70, Syk, Shc, PI3 kinase, and Grb2. Since CLASPs have an extracellular domain, CLASPs protein can independently initiate a signal transduction cascade through engagement of its extracellular domain. Otherwise CLASPs may cooperate with an antigen receptor signaling complex (*e.g.*, with CD3/TCR, BCR, FcR), to facilitate tyrosine kinase signal transduction

The ITAMs have demonstrated different binding specificity and affinities for SH2 domains (Clements, *et al.*, 1999, Ann. Rev. Immunol. 17: 89-108). For example, Shc, PI3 kinase, and Grb2 bind to dual and mono phosphorylated ITAMs with different affinities. Thus the ITAMs in CLASPs are believed to provide quantitative as well as qualitative differences in signal transduction depending up their phosphorylation state, as well as to inhibit or augment specific protein interactions and hence specific tyrosine kinase-mediated signaling pathways in leukocytes.

Antagonizing the PTK-CLASP-2 interaction (*e.g.*, phosphorylation of CLASP-2) will thus inhibit immune function. In one embodiment, interactions between ITAM-bearing human CLASPs and their binding partners are believed to be antagonized by the alpha subtype (SIRPalpha) of signal regulatory proteins that has been shown to negatively

regulate ITAM-dependent lymphocyte activation (Lienard H; 1999, J Biol Chem 274: 32493-9). Also, a recently recognized family of immunoreceptor tyrosine-based inhibition motif (ITIM) receptors are thought to inhibit the ITAM-induced activation of immune competent cells (Gergely, *et al.*, 1999, J. Immunol Lett 68: 3-15) and therefore may block CLASP-partner interaction.

#### **5.2.1.6. DOCK**

CLASP-2 polypeptides contain a new “DOCK” motif, not previously described in the scientific literature. The CLASP DOCK motif includes a series of five tyrosines surrounded by conserved sequences in regions A, B, C, D, and G (see FIG. 5B).

There are also two highly conserved non-tyrosine containing regions (E and G) separated by nine amino acids (P+EXAI+XM) and (LXMXL+GXVXXXVNXG) (where X is any amino acid).

The cytoplasmic region of CLASP-2 immediately following the ITAM domains exhibits sequence similarity to the C-terminal third of the so-called “DOCK” proteins. The DOCK gene family includes three molecules that are the human homologues of the *C. elegans* CED proteins known to be involved in apoptosis. CED-5 (DOCK180), a major CRK-binding protein, alters cell morphology upon translocation to the membrane (mediates the membrane motion that scavenger cells exhibit as they surround and engulf dying cells; its function can be partially rescued by the human DOCK180 (Wu *et al.*, 1998, *Nature* 392: 501-504). Myoblast City in *Drosophila* (MBC) is another member of the DOCK protein family and has been found to be involved in myoblast fusion (Erickson, *et al.*, 1997, *J. Cell Biol.* 138: 589). Since CLASP-2 expression is found in syncytial tissues such as placenta, muscle, and heart, it is believed that CLASP-2 is involved in mediating or inhibiting cell fusion.

The DOCK family has been implicated in the control of cell shape. DOCK1, when transfected into spindle cells, can make them flattened and polygonal (Takai, *et al.*, 1996, *Genomics* 35: 403-303). DOCK1 expression is ubiquitous except in hematopoietic cells. DOCK2 is expressed in hematopoietic cells and when transfected into spindle cells can make them round up (Nishihara, H., 1999, *Hokkaido Igaku Zasshi* 74: 157-66). DOCK2 is expressed in peripheral blood lymphocytes, thymus, spleen, and liver.

#### **5.2.1.7. COILED-COIL**

CLASP-2s have the two coiled-coil domains (Lupas *et al.*, 1991, *Science* 252: 1162-64; Lupas, A., 1996, *Meth. Enzymology* 266: 513-525). Coiled-coil domains are known to interact directly with cytoskeleton, indicating that CLASP-2 proteins interact directly with the cytoskeleton. Thus, it is believed that CLASP-2 binds cytoskeletal proteins, *e.g.*, spectrin, ankyrin, hsp70, talin, ezrin, tropomyosin, myosin, plectin, syndecans, paralemmin, Band 3 protein, Cytoskeletal protein 4.1, Tyrosine phosphatase PTP36 and other molecules.

#### **5.2.1.8. PDZ Ligand**

CLASP-2 proteins contain a PDZ-ligand motif ("PBM" or "PL") at the C-terminus of the protein. This short (3 – 8 amino acid) motif mediates the binding of proteins terminating at their carboxyl terminus in the motif (most commonly S/T – X – V – free carboxyl-terminus) to other proteins containing one or more specific PDZ domains (See Songyang *et al.*, 1997, *Science* 275: 72 and Doyle *et al.*, 1996, *Cell* 85: 1067 for a discussion of PDZ-ligand structures).

PDZ domain-containing proteins are involved in the organization of ion channels and receptors at the neurological synapse and in establishing and maintaining polarity in epithelial cells via their binding to the C-termini of transmembrane receptors. It has been shown that PDZ-domain containing proteins can mediate protein-protein interactions in immune system cells (*e.g.*, DLG1 binds to the lymphocyte potassium channel KV1.3 in human T lymphocytes, (Hanada *et al.*, 1997, *J. Biol. Chem.* 272: 26899).

Biochemical evidence that CLASP-2 interacts with the PDZ domains of three closely related proteins is shown in FIG 9A-D. FIG. 9A demonstrates the specificity of the interaction, as the C-terminal 20 amino acids of CLASP-2 bind PSD-95, NeDLG, and DLG1, but not to the PDZ domains of the TIAM-1 protein. FIG. 9B demonstrates the affinity of the interaction. Notably, the highest affinity interaction occurs between CLASP-2 and NeDLG, with a specific binding affinity of at least  $10^4 \text{ M}^{-1}$ . Affinities in the micromolar range have been found for other biologically important PDZ-ligand interactions. FIG. 9C demonstrates the ability to inhibit CLASP-2 PDZ interactions using either a short fragment of CLASP-2 (the eight C-terminal amino acids) or the C-terminus of KV1.3. As noted above, KV1.3 is known to bind to DLG1 in live lymphocytes. FIG. 9D demonstrates that CLASP-2 and KV1.3 compete for PDZ binding; *i.e.*, not only does KV1.3 block CLASP-2 binding but



CLASP-2 also blocks KV1.3 binding. The ability of the eight C-terminal residues of CLASP-2 to inhibit the interaction of both CLASP-2 and KV1.3 with selected PDZ domains suggests that compounds related to the C-terminal eight-amino acids of CLASP-2, when introduced into cells, will mediate changes in multiple protein-protein interactions involved in the function of lymphoid tissues and other tissues that express these proteins (including heart, lung, and kidney).

Evidence that the C-terminal 8 amino acids of CLASP-2, when introduced into cells, can effect cellular function comes from the experiments in which these amino acids were introduced into cells as a fusion, *e.g.*, with the HIV-derived TAT transporter peptide sequence. Addition of the TAT-CLASP-2 fusion peptide to Jurkat T lymphocytes (compared to controls using the TAT peptide alone) results in subtle, time-dependent alterations in intracellular calcium concentrations as measured using the calcium indicator dye Fluo-4. While these results are consistent with the hypothesis that the TAT-CLASP-2 fusion changes T cell ion fluxes. In particular, the results indicate that the CLASP-2 C-terminal sequence can slightly increase basal intracellular calcium concentrations and can slightly decrease the proportional increase in calcium upon activation of the cells with anti-CD3 antibody. Such changes would be expected for a compound that disrupts localization of the T cell activation-associated CLASP-2 protein and the KV1.3 potassium channel. Small changes in T cell calcium flux can result in large changes in the functional activity of the cells (Wulfing *et al.*, 1997, J. Exp. Med. 185: 1815).

#### **5.2.1.9. Modulation of Immune Responses**

CLASP-2 proteins, as described above, modulate immune function in a variety of ways and through a variety of mechanisms (*i.e.*, changing the threshold for T cell activation) by affecting formation of the immunological synapse. Establishment and maintenance of the immunological synapse can involve: (A) signal transduction, (B) cell-cell interactions, and (C) membrane organization.

##### **(A) Signal transduction**

Human CLASP proteins, as discussed above, contain SH3 domains and tyrosine phosphorylation sites. These regions have been shown to be involved in signal transduction in a variety of cells including lymphocytes. Thus, human CLASP proteins are believed to interact with these regions during signal transduction events which lead to modulation of immune responses.

CLASP proteins can interact with Tec sub-family of nonreceptor tyrosine kinases. The Tec sub-family of nonreceptor tyrosine kinases consists of Tec, Btk, Tsk/Itk/Emt Itk, and Bmx, and is defined by the presence of SH3 and SH2 domains adjacent to the catalytic domain and an amino-terminal region containing a pleckstrin homology (PH) domain, a Tec homology (TH) domain, and a proline-rich region (Mano, H.; 1999, Cytokine Growth Factor Rev 10: 267-80). The T cell specific Tsk/Itk/Emt, and Btk expressed in most hematopoietic cells other than T cells are important components of antigen receptor signaling pathways in hematopoietic cells.

Btk has been identified as the gene defective in murine X-linked immunodeficiency (*xid*) and human X-linked agammaglobulinemia (XLA) (Nisitani, S., 2000, Proc Natl Acad Sci U.S.A. 97: 2737-42). In *xid* mice, B cell numbers are reduced to one-half of normal and the titers of specific immunoglobulin isotypes are significantly reduced; in addition, *xid* B cells are insensitive to a number of mitogenic stimuli. The human disorder is much more severe, resulting in nearly complete elimination of the B cell compartment and dramatically reduced immunoglobulin levels. Biochemical studies have supported multiple roles for Btk in B cell activation. Btk kinase activity and tyrosine phosphorylation are increased after cross-linking either the B cell receptor on B cells or the high affinity IgE receptor, FcRI, on mast cells. Interleukin-5 and interleukin-6 treatment have also been shown to lead to the activation of Btk.

Itk, like Btk, is tyrosine-phosphorylated upon antigen receptor cross-linking (Mano, H., 1999, Cytokine Growth Factor Rev, 10: 267-80). In addition, peripheral T cells from mice lacking functional Itk are refractory to stimulation by antibodies to CD3 plus antigen presenting cells. These Itk-deficient T cells can be stimulated by phorbol ester and calcium ionophore, demonstrating that Itk acts in signaling pathways proximal to the TCR.

Unlike the related Src family tyrosine kinases including Lyn, Lck, Fyn, ZAP-70, SyK, and CSK, the Tec family kinases lack the amino-terminal myristylation site crucial for the membrane localization of Src family kinases, suggesting that some adaptor proteins are required for the their membrane localization (Mano, H., 1999, Cytokine Growth Factor Rev 10: 267-80). Since all the Tec family kinases contain a proline-rich region which could be bound by a SH3 domain, and since all the human CLASPs contain a SH3 domain, it is believed that human CLASPs could serve as adaptors for the members in the Tec family in different hematopoietic cells.

GTP-binding proteins play an important role in immune response (Mach, B., 1999, Science 285: 1367). A number of biochemical events triggered by TCR/CD3-induced



T cell activation are ablated by agents that modulate the action of G proteins. Pertinent to this is the ability of cholera toxin to inhibit the cellular proliferation and intracellular  $\text{Ca}^{2+}$  mobilization that is mediated by anti-CD3 antibody treatment of T cells. The G protein competitive inhibitor GDPS, can impede the extent of inositol phosphates generated upon stimulation in peripheral T lymphocytes. Nonhydrolyzable analogs of GTP, such as GTPS, or other agents such as ALF that activate G proteins by circumventing the need for receptor engagement, can result in T cell activation.

The Gαq/11 subfamily (Stanners, J., 1995, J Biol Chem 270: 30635-42) and Rap1 (Lafont, V., 1998, Biochem Pharmacol 55: 319-24) of GTP-binding proteins have been shown to be involved in human T cell receptor/CD3-mediated signal transduction pathway. Also, Cdc42, a Rho family small GTPase, is known to play a critical role in the formation of actin microspikes in response to external stimuli (Miki, H.; 1998, Nature, 391: 93-6). Interestingly, a Cdc42 binding protein, WASP, has a proline-rich domain which could interact with the SH3 domain present in all the human CLASPs. Human CLASPs may interact with these GTP-binding proteins.

Several adaptor proteins including NCK, CBL (Bachmaier, K., 2000 Nature 403: 211-6), SHC, LNK, SLP-76, HS1, SIT, VAV, GrB2, and BRDG1, and two tyrosine phosphatases, EZRIN, SHP-1 and SHP-2 have been shown to interact with ITAM or SH3 domains. These proteins may also interact with CLASP-2. Several proteins have been shown to interact with ITAM or SH3 domains and may also interact with CLASP-2. These include adaptor proteins such as NCK, CBL (Bachmaier, K., 2000, Nature 403: 211-6), SHC, LAT, LNK, SLP-76 (Krause M *et al.*, 2000, J Cell Biol 149: 181-94), HS1, SIT, VAV, GrB2 (Zhang W. and Samelson, L.E., 2000, Semin Immunol 12: 35-41), and BRDG1, kinases such as SYK and LCK, and tyrosine phosphatases such as SHP-1 and SHP-2. These interactions can be defined by a number of different biochemical or cell biological methods including in vitro binding assays, co-immunoprecipitation assays, co-immunostaining (Harlow, E. and Lane, D., 1999, Using Antibodies: A laboratory Manual. Cold Spring Harbor Press) or genetic assays such as yeast the yeast two hybrid system, in which a CLASP-2 protein or fragment can be used as “bait” (Zervos *et al.*, 1993, Cell 72: 223-232; Madura *et al.*, 1993, J. Biol. Chem 268: 12046-12054).

Other assays include in vitro binding assays, co-immunoprecipitation assays, co-immunostaining assays, and yeast two hybrid system screening assays in which a CLASP-2 domain or fragment can be used as “bait” or “trap” protein (Zervos *et al.* (1993), Cell 72: 223-232; Madura *et al.* (1993) J. Biol. Chem. 268: 12046-12054).

In other embodiments, CLASP polypeptides are transfected into lymphocytes. After transfection, a variety of standard assays can be used to evaluate, for example, CLASP modulation of T cell activation. These assays include calcium influx assays, NF-AT nuclear translocation assays (*e.g.*, Cell, 1998, 93: 851-61), NF-AT/luciferase reporter assays (*e.g.*, MCB 1996 16: 7151-7160), tyrosine phosphorylation of early response proteins such as HS1, PLC- $\gamma$ , ZAP-76, and Vav (*e.g.*, J. Biol. Chem. 1997, 272: 14562-14570).

#### (B) Cell-Cell Interaction

As discussed above, human CLASP proteins are homologues of E-cadherin. As shown in FIG. 1, CLASP-2 contains both a cadherin cleavage domain and a cadherin ectodomain. Therefore CLASP-2 proteins may interact with cadherins through these domains. The cadherins constitute a family of cell surface adhesion molecules that are involved in calcium-dependent cell to cell adhesion. Human cadherins, E-, P- N- and VE-cadherin, have a restricted tissue distribution: E- and P-cadherin are expressed in epithelial tissues, N-cadherin is found mainly on neural cells, and VE-cadherin is found on vascular endothelium. Homophilic binding between cadherins on adjacent cells is vital for the maintenance of strong cell to cell adhesion in these tissues. For example E-cadherin is required for the formation of adherens junctions between mature epithelial cells and is involved in Langerhans cell adhesion to keratinocytes, and VE-cadherin is needed for the maintenance of lateral association between endothelial cells. The extracellular regions of mature mammalian cadherins are comprised of five "CAD" modules of approximately 1110 amino acids. Crystallographic and biochemical studies indicate that cadherins can form dimers on the cell surface, and that interaction with dimeric cadherins on opposing cell surfaces can lead to the formation of "zipper-like" cell junctions.

The integrins are a second family of transmembrane adhesion molecules that are involved in both cell to cell and cell to matrix interactions. At least 15 chains associate with 8 chains to form a large number of heterodimeric integrins that can be classified into several major subfamilies based on their shared use of a particular chain. Members of three subfamilies, the 1, 2, and 7 integrins, are commonly found on leukocytes. The expression of 1 integrins is widespread (for example, 51, CD49e/CD29, is found on T cells, granulocytes, platelets, fibroblasts, endothelium, and epithelium), whereas the 2 and 7 integrins have a restricted pattern of expression.

Interestingly, E-cadherin on human epithelial cells has been found to be a ligand for the mucosal lymphocyte integrin, E7, and a similar interaction has been indicated

in the mouse. Monoclonal antibodies to E-cadherin or to E7 block IEL adherence to epithelial cells, and transfection of cells with E7 confers upon them the ability to adhere to cells transfected with E-cadherin.

L929 cells can be transfected with CLASP-2 and Neomycin. G418-resistant clones can be screened for CLASP-expression with anti-CLASP peptide-specific antibodies. CLASP-expressing clones can be used to test for homotypic and/or heterotypic calcium dependent cell adhesion using the "cell aggregation assay" described for cadherin molecules (Murphy-Erdosh, C. *et al.*, 1995, J. Cell Biol. 129: 1379-1390).

Several approaches can be used to identify the amino acids involved in the binding domains. Soluble fusion molecules (*e.g.*, EC12-IgG, ECC-IgG, ECM-IgG, and GST-EC12), peptides, and peptide-specific anti-CLASP antibodies are available for blocking experiments in the above-described assay. Transfectants generated by site-directed mutagenesis can also be used.

#### (C) Membrane Anchoring/Cytoskeletal Interactions

Interestingly, tyrosine-phosphorylated ITAMs interact with actin cytoskeleton upon activation of mature T lymphocytes (Rozdzial, M. M., 1995, Immunity 3: 623-633). Since human CLASPs contain both ITAMs and coiled-coil domains which have been shown to interact with cytoskeletal proteins, CLASPs are believed to play an important role in modulating cell surface molecule expression by re-organizing cytoskeletal structure.

F-actin microfilament cytoskeletal organization has been known to be involved in the modulation of cell surface molecule expression. WASP, a GTPase-binding protein, plays a critical role in the formation of actin microspikes in response to external stimuli and ectopic expression of WASP induces the formation of F-actin filament clusters that overlap with the expressed WASP itself. Another WASP family protein, N-WASP, has also been shown to play important roles in filopodium formation. Both of these proteins cause actin polymerization, but with different features when they are expressed in cells; WASP mainly localizes at perinuclear areas and causes actin clustering, but most N-WASP is present at plasma membranes and induces filopodium formation (Miki, H.; 1998, Nature 391: 93-6). Both WASP and N-WASP, contain a proline-rich domain which could interact with the SH3 domain present in all the human CLASPs. CLASP-2 may interact with F-actin filament through CLASP-2 binding to WASP or WASP-like proteins.

Standard assays can be used for detecting CLASP protein interaction with cytoskeletal proteins. These assays include co-sedimentation assays, far western blot analysis (Ohba, T., 1998, Anal. Biochem. 262: 185-192), surface plasmon resonance, F-actin staining

with phalloidin in CLASP-transfected lymphocytes (*e.g.*, Small, J. *et al.* 1999, Microsc. Res. Tech. 4: 3-17), and immunocytochemical analysis of subcellular distribution of focal adhesion proteins (such as paxillin, tensin, vinculin, talin, and FAK in CLASP-transfected lymphocytes; see, *e.g.*, Ridyard, M.S., 1998, Biochem. Cell Biol. 76: 45-58).

### 5 **5.2.2. CLASP-2 Exon Structure and Genomic Domains**

Alternative splicing is likely to represent a regulatory switch that governs different functions of CLASP-2 in immune responses. Additionally, alternative splice variants affecting the untranslated regions of an RNA can be a way of regulating RNA stability.

As noted *supra*, CLASP-2 gene expression is characterized by alternative exon usage. Intron/exon structure can be predicted by computer analysis of genomic DNA, however, splice junctions and alternative splicing can only be elucidated by comparison of genomic clones to cDNA clones. Alternative splicing and RNA editing are mechanisms generate a variety of proteins from the same gene. An example for how alternative splicing is used to generate thousands of different proteins from only a few genes is represented by the Neurexin gene family (for review of Neurexins, see Missler M. and Suedhof, T., 1998, Trends in Genetics, 14: 20-25). Comparative analysis of CLASP-2 genomic clones and cDNA clones revealed that CLASP-2 is composed of numerous exons and that distinct CLASP-2 transcripts are generated by alternative splicing. The protein encoding portion of CLASP-2 is covered by at least 14 exons (FIG. 6A).

Numerous diseases are caused or are thought to be caused by splice site mutations that can cause exon skipping or otherwise result in a truncated protein product. Some of these diseases include, *e.g.*, Marfan Syndrome (Liu W, *et al.*, 1997, Nat. Genet. 16: 328-9), Hunter disease (Bonucelli G, *et al.*, 2000, Hum. Mutat. (Online) 2000 15(4): 389, Duchenne muscular dystrophy (Wibawa T, *et al.*, 2000, Brain Dev. 22(2): 107-112), Myelomonocytic leukemia (Wutz D, *et al.*, 1999, Leuk. Lymphoma 35: 491-9.), and Isovaleric acidemia (Vockley J, *et al.*, 2000, Am. J. Hum. Genet. 66: 356-67). This is especially true for genes composed of many exons (such as CLASP-2). The genomic sequence around CLASP-2 exon/intron boundaries is useful for diagnostic approaches towards the identification of diseases caused by splice site mutations. The abundance or presence of CLASP-2 isoforms in cell populations (*e.g.*, hematopoietic cells, lymphocytes) is correlated with a disease state by comparing the abundance of CLASP-2 in cells from

subjects suffering from the disease with the level of CLASP-2 in cells from healthy subjects. This can be accomplished by utilizing any number of assays (e.g., PCR).

Alignment of the CLASP-2 intron/exon splice sites with the CLASP-2 protein sequence and the finding of conserved exon/intron boundaries within the CLASP gene family (FIG. 6) suggest that specific CLASP-2 exons encode functionally distinct protein domains (see FIG. 6 and Example 4). ITAM and DOCK motifs 1 and 2 are encompassed by splice sites (amino acid residues 946 and 1063); DOCK motif 3 and COILED-COIL motif 1 and 2 are also encompassed by splice sites (amino acid residues 1102, 1170 and 1246, respectively).

CLASP-2 alternative transcripts are summarized in FIG. 3 and FIG.11B. Briefly, one alternative exon missing in CLASP-2A is present in CLASP-2B and CLASP-2D. This exon contains the DNA portion encoding the ITAM motif and DOCK motif 1. The CLASP-2D protein product does not contain the C-terminal 38 amino acids of CLASP-2A and CLASP-2B. Thus, a PDZ binding motif (SSVV; amino acid residue 1286 through 1289) that is only present in the CLASP-2A/B-specific C-terminal end is missing in the CLASP-2D gene product. The presence or absence of this PDZ binding motif can be attributed to alternative RNA processing. Additionally, a CLASP-2 alternative transcript has been found that deletes nucleotides 209-291 that results in a premature stop codon. The protein encoded by this transcript appears to be a soluble form of CLASP-2 that may regulate (e.g., is an antagonist or an agonist) the function other CLASP family members and isoforms.

### **5.2.3. CLASP Superfamily Members**

As is illustrated in FIG. 5, CLASP-2 is a member of a superfamily of immune-cell associated proteins with similar motifs. CLASP-1 was described in U.S.S.N. 09/411,328, filed October 1, 1999. CLASP-1 uniquely among the known CLASPs contains SH3 binding domain motifs. CLASP-2A, -B, -C, and -E polypeptides have no adaptor binding sites or SH3 binding domains found in CLASP-1. CLASP-3, CLASP-4, CLASP-5 and CLASP-7 are described in copending U.S.S.N. 60/182,296, filed February 14, 2000, and which is incorporated by reference herein in its entirety for all purposes.

### **5.3. CLASP-2 mRNA Expression**

As described in Example 2, CLASP-2 mRNA expression was assayed in tissues and cell lines by Northern analysis. The results are shown in FIG. 4A and B. The



results of Northern Analysis of CLASP-2 expression and expression of other members of the CLASP family are summarized in Table 2.

**Table 2**

Tissue/Cell Line <sup>1</sup>	CLASP					
	1	2 <sup>3,4</sup>	3	4	5	7
PBL	+ <sup>2</sup>	-	-	++++	++	-
Lung	-	+	-	-	-/+	+++
Placenta	-/+	+++	+	-/+	+	+
Sm Intestine	-/+	-	-	-	-/+	+
Liver	-/+	-/+	-/+	-	-/+	+
Kidney	-/+	+	+++	-/+	+	++
Spleen	++	-	-	-/+	+	-/+
Thymus	++	-	-	-/+	+	-
Colon	-	-	-	-	-	-
Skel Muscle	-	-/+	++	-	-	-/+
Heart	-/+	++	+++	-/+	-	+++
Brain	+++	-/+	-/+	-	-	-
Jurkat	++	++	++	+	-	-
MV411	++	-	++	+	+	+
THP1	++	-	-	-	-	-/+
HL60	-	-	-	-	-/+	-
9D10	++	++ <sup>5</sup>	+	+	+	+
3A9	+	-/+	-	-	-	-
CH27	+	-/+	-	-	-	-
293	-	++	+++	+	-	+

1. Jurkat = human T cell line; MV4-11 = B myelomonocyte; 9D10 = B cell line; THP-1 = monocyte; 3A9 = mouse T cell; CH27 = mouse B cell line; HL60 = human promyelocyte; 293 = embryonic kidney epithelial cells (293)

2. Table Legend (based on Northern blot results): - = no expression; -/+ = low expression; + = medium expression; ++ medium high expression; +++ high expression.

3. A CLASP-2 EST (EST 815795) was identified from a bone marrow cDNA library.

4. The probe used (HC2.2) did not distinguish between CLASP-2A, -2B, -2C and 2D.. This probe encompasses nucleotides 3920 to 4650 (731 bp long) from CLASP-2A cDNA.

5. In RNA from 9D10, the major transcript runs substantially shorter than the major transcripts seen in Jurkat and 293 cells; however, the longer transcript is also present in 9D10. Hybridization of probe HC2.2 with 9D10 total RNA reveals at least 3 different transcripts. See FIG. 4B

As indicated in Table 2 and shown in FIG. 4, CLASP-2 is expressed most strongly in placenta followed by lung, kidney and heart; CLASP-3 is expressed strongly in kidney and heart, and less strongly in placenta and skeletal muscle ; CLASP-4 is expressed exclusively in peripheral blood lymphocytes; CLASP-5 is expressed strongly in peripheral blood leukocytes, present in placenta, kidney, spleen and thymus, and weakly in lung, small

intestine and liver. It is not expressed in brain, heart, skeletal muscle and large intestine; CLASP-7 is expressed strongly in lung, heart, liver and kidney, but not in PBL, brain or thymus.

Differences in tissue expression patterns for different CLASP proteins indicate different CLASPs have differential roles in immune function and, accordingly, can be separately targeted to achieve different functions. For example, since CLASP proteins are necessary for proper function or signaling by the T cell receptor (TCR), the tissue specific distribution of different CLASPs permits differential modulation of the immune response in different tissues. Since CLASP-2 is present in heart, blocking CLASP-2 function or expression is useful to selectively block immune response in the heart (for example, to selectively stop immune response in the heart compartment, *e.g.*, following cardiac transplant rejection or post-MI inflammation, without compromising immunity elsewhere. Similarly, blocking CLASP-3 can block rejection of the kidney following kidney transplant. Furthermore, by adjusting the level of inhibition, the degree of immune blockage versus response can be modulated in the compartments represented by each CLASP.

#### **5.4. CLASP-2 Polynucleotides And Methods Of Use**

The present invention provides a variety of CLASP-2 polynucleotides and methods for using them. In one aspect, the polynucleotide of the invention encodes a polypeptide comprising at least a fragment (*e.g.*, an immunogenic fragment) of a CLASP-2 protein (*e.g.*, at least a fragment of SEQ. ID. NO: 2, 4, 6 or 10) or variant thereof. In another aspect, the molecules that comprise a CLASP-2 polynucleotide that, while not necessarily encoding a CLASP-2 protein or fragment, is useful as a probe or primer for detecting CLASP-2 expression, for inhibition of CLASP-2 expression (*e.g.*, antisense or ribozyme-mediated inhibition), for gene knockout, and the like.

##### **5.4.1. CLASP-2 Polynucleotides**

The invention also provides isolated or purified nucleic acids having at least 8 nucleotides (*i.e.*, a hybridizable portion) of a CLASP-2 sequence or its complement; in other embodiments, the nucleic acids consist of at least about 25 (continuous) nucleotides, about 50 nucleotides, about 100 nucleotides, about 150 nucleotides, about 200 nucleotides, about 250 nucleotides, about 500 nucleotides, about 550 nucleotides, about 600 nucleotides, or about 650 nucleotides or more of a CLASP-2 sequence, or a full-length CLASP-2 coding sequence. In another embodiment, the nucleic acids are smaller than about 35, about 200 or about 500

nucleotides in length. Polynucleotides can be single or double stranded, and may be DNA, RNA, PNA or a hybrid molecule.

In specific aspects, nucleic acids are provided which comprise a sequence complementary to at least about 10, 25, 50, 100, 150, 200, 250, 500, 550, 600, or 650 nucleotides or the entire coding region of a CLASP-2 coding sequence. Usually, the isolated polynucleotide is less than about 100 kbp, generally less than about 50 kbp, and often less than about 20 kbp, less than about 10 kbp, less than about 5 kbp, or less than about 1000 nucleotides in length.

In a specific embodiment, a nucleic acid that is hybridizable to a CLASP-2 nucleic acid or its complement, or to a nucleic acid encoding a CLASP-2 derivative, under conditions of low stringency is provided. Derivatives of CLASP-2 contemplated include, but are not limited to, splice variants of a gene encoding a CLASP-2, other members of a CLASP-2 gene family which differ from one of the CLASP-2 nucleotide or amino acid sequences disclosed herein by the insertion or deletion of one or several domains, and the like.

In one embodiment, the CLASP-2 polynucleotide is identical or exactly complementary to SEQ. ID NO: 1, 3, 5 or 9 or selectively hybridizes to an aforementioned sequence. In various embodiments, the polynucleotide is identical or exactly complementary to, or selectively hybridizes to, the nucleotide sequence encoding a particular protein domain or region, or a particular gene exon of the CLASP-2 mRNA or genomic sequence. Such polynucleotides are particularly useful as probes, because they can be selected to identify a defined species of CLASP-2.

In addition to the polypeptide and polynucleotide sequences specifically exemplified herein, the invention contemplates CLASP-2 homologues from other species, allelic and splice variants, and other variants disclosed herein. The CLASP-2 gene exhibits evidence of alternative splicing of transcripts.

For example, CLASP-2A and CLASP-2C are related to each other as apparent splice variants, with CLASP-2C containing an exon not found in CLASP-2A. The exon sequence is 5'-AGG GAT TTT GAG AGG CTG GCC CAT CTG TAT GAC ACG CTG CAC CGG GCC TAC AGC AAA GTG ACC GAG GTC ATG CAC TCG GGC CGC AGT TNC TGG GGA CCT ACT TCC GGG TAG CCT TCT TCG GGC AG-3' (encoding the peptide sequence: RDFERLAHLYDTLHRAYSKVTEVMHSGRRLGTYFRVAFFGQGF). It will be apparent to one of skill that, by using polynucleotide probes or primers corresponding to the nucleic acid sequence above, or by using antibodies that specifically

recognize the peptide above, or those polynucleotide probes or primers shown in Table 3 below, it is possible to distinguish between different CLASP isoforms(*e.g.*, to detect differential expression).

**Table 3**

	Found in/will detect	Exemplary Probe/Primer (5' – 3')	Notes/Comments
1	full length hC2A	F1: CCCAGATTTTATGATGAG R1: GATAATGACAAAGTTCTGAC	
2	full length hC2D	F2: CTGGAAATCTTGACAAAAATGC R2: GTCCTTTTAATACAGATGTGG	
3	hC2B, hC2C, hC2E	F3: GAGAGGCTGGCCCATCTGTATG R3: ATCTTCAAAGAATCCCTGCC	Distinction based upon product size differences following PCR
4	hC2D	F4: GAAGCAGTCCAGTGGGAGCCG R4: GCCTCCCCGGCTCCTCCTCAGG	Recognizes hC2D-specific insertion
5	hC2D	F3: GAGAGGCTGGCCCATCTGTATG R5: CCTCCACATCTGTTTCACTGTC	
6	hC2E	F5: CTCATGATGGAAGACGTGGG R6: GATGAGCTCGTAGCGCTCGGC	Spans deletion unique to hC2E. Distinction based upon product size differences following PCR
7	hC2B	F6: CATTGGCGTTTAAGCTCCTG R3: ATCTTCAAAGAATCCCTGCC	F6 primer spans deletion unique to hC2E
8	hC2A	F7: GGACCCATAGTTCATGATCG R4: CTTATCTTCAAGAAATCCCTC	R4 primer spans the region where other CLASPs have an insert

#### **5.4.1.1. Substantial Identity**

In some embodiments, the CLASP-2 polynucleotides of the invention are substantially identical to SEQ ID NOs: 1, 3, 5, or 9, or to a fragment thereof.

An indication that two nucleic acid sequences are substantially identical is that the two polynucleotides have a specified percentage sequence identity *e.g.*, usually at least about 75%, at least about 80%, at least about 85%, at least about 90%, at least about 95%, or at least about 98 identity over a specified region when optimally aligned.

Another indication that two nucleic acid sequences are substantially identical is that a polypeptide encoded by the first nucleic acid is immunologically cross reactive with the antibodies raised against the polypeptide encoded by the second nucleic acid, as described below. Thus, a polypeptide is typically substantially identical to a second polypeptide, for example, where the two peptides differ only by conservative substitutions. Another indication

that two nucleic acid sequences are substantially identical is that the two molecules or their complements hybridize to each other under stringent conditions, as described below.

Yet another indication that two nucleic acid sequences are substantially identical (*e.g.*, a naturally occurring allele of the CLASP-2 sequence of SEQ ID NO: 1) is that the same primers can be used to amplify the sequence. For example, CLASP-2 polynucleotides can be PCR amplified from cDNA derived from human lymphocytes using the primer pairs shown in Table 3.

The primers of Table 3 are also useful for amplification of CLASP-2 splice variants. Another indication that two nucleic acid sequences are substantially identical is that they selective hybridize under stringent conditions (*i.e.*, one sequence hybridizes to the complement of the second sequence), as described *infra*.

#### **5.4.1.2. Selective Hybridization**

The invention also relates to nucleic acids that selectively hybridize to exemplified CLASP-2 sequences (including hybridizing to the exact complements of these sequences). Selective hybridization can occur under conditions of high stringency (also called “stringent hybridization conditions”), moderate stringency, or low stringency.

##### **5.4.1.2.1. High Stringency**

“Stringent hybridization conditions” are conditions under which a probe will hybridize to its target subsequence, typically in a complex mixture of nucleic acid, but not to other sequences. Stringent conditions are sequence-dependent and will be different in different circumstances. Longer sequences hybridize specifically at higher temperatures. An extensive guide to the hybridization of nucleic acids is found in Tijssen, Techniques in Biochemistry and Molecular Biology--Hybridization with Nucleic Probes, “Overview of principles of hybridization and the strategy of nucleic acid assays” (1993). Generally, stringent conditions are selected to be about 5-10°C lower than the thermal melting point ( $T_m$ ) for the specific sequence at a defined ionic strength pH. The  $T_m$  is the temperature (under defined ionic strength, pH, and nucleic concentration) at which 50% of the probes complementary to the target hybridize to the target sequence at equilibrium (as the target sequences are present in excess, at  $T_m$ , 50% of the probes are occupied at equilibrium). Stringent conditions will be those in which the salt concentration is less than about 1.0 M sodium ion, typically about 0.01 to 1.0 M sodium ion concentration (or other salts) at pH 7.0 to 8.3 and the temperature is at least about 30°C for short probes (*e.g.*, 10 to 50 nucleotides)



and at least about 60°C for long probes (e.g., greater than 50 nucleotides). Stringent conditions may also be achieved with the addition of destabilizing agents such as formamide. For high stringency hybridization, a positive signal is at least two times background, preferably 10 times background hybridization. Exemplary high stringency or stringent hybridization conditions include: 50% formamide, 5x SSC and 1% SDS incubated at 42° C or 5x SSC and 1% SDS incubated at 65° C, with a wash in 0.2x SSC and 0.1% SDS at 65° C. In a specific embodiment, a nucleic acid which is hybridizable to a CLASP-2 nucleic acid under the following conditions of high stringency is provided: Prehybridization of filters containing DNA is carried out for 8 h to overnight at 65°C in buffer composed of 6X SSC, 50 mM Tris-HCl (pH 7.5), 1 mM EDTA, 0.02% PVP, 0.02% Ficoll, 0.02% BSA, and 500 µg/ml denatured salmon sperm DNA. Filters are hybridized for 8-16 h at 65°C in prehybridization mixture containing 100 µg/ml denatured salmon sperm DNA and 5-20 X 10<sup>6</sup> cpm of <sup>32</sup>P-labeled probe. Washing of filters is done at 65°C for 15-30 h in a solution containing 2X SSC, 0.1% SDS. This is followed by a wash in 0.2X SSC and 0.1% at 50°C for 15-30 min before autoradiography.

#### **5.4.1.2.2. Moderate Stringency**

In another specific embodiment, a nucleic acid, which is hybridizable to a CLASP-2 nucleic acid under conditions of moderate stringency is provided. Examples of procedures using such conditions of moderate stringency are as follows: Filters containing DNA are pretreated for 6 h at 55°C in a solution containing 6X SSC, 5X Denhart's solution, 0.5% SDS and 100 µg/ml denatured salmon sperm DNA. Hybridizations are carried out in the same solution and 5-20 X 10<sup>6</sup> cpm <sup>32</sup>P-labeled probe is used. Filters are incubated in hybridization mixture for 12-16 h at 55°C, and then washed twice for 30 minutes at 50°C in a solution containing 1X SSC and 0.1% SDS. Filters are blotted dry and exposed for autoradiography. Other conditions of moderate stringency which can be used are well-known in the art. Washing of filters is done at 45°C for 1 h in a solution containing 0.2X SSC and 0.1% SDS.

#### **5.4.1.2.3. Low Stringency**

By way of example and not limitation, procedures using such conditions of low stringency are as follows (see also Shilo and Weinberg, 1981, Proc. Natl. Acad. Sci. U.S.A. 78: 6789-6792): Filters containing DNA are pretreated for 6 h at 40 C in a solution

containing 35% formamide, 5X SSC, 50 mM Tris-HCl (pH 7.5), 5 mM EDTA, 0.1% PVP, 0.1% Ficoll, 1% BSA, and 500 µg/ml denatured salmon sperm DNA. Hybridizations are carried out in the same solution with the following modifications: 0.02% PVP, 0.02% Ficoll, 0.2% BSA, 100 g/ml salmon sperm DNA, 10% (wt/vol) dextran sulfate, and 5-20 X 10<sup>6</sup> cpm <sup>32</sup>P-labeled probe is used. Filters are incubated in hybridization mixture for 18-20 h at 40 C, and then washed for 1.5 h at 55 C in a solution containing 2X SSC and 0.1% SDS. The wash solution is replaced with fresh solution and incubated an additional 30 minutes at 50-55°C. Filters are blotted dry and exposed for autoradiography. If necessary, filters are washed for a third time at 60-65°C and reexposed to film. Other conditions of low stringency that can be used are well known in the art (e.g., as employed for cross-species hybridizations).

#### **5.4.1.3. CLASP-2 Variants and Fragments**

The CLASP-2 variants of the invention can contain alterations in the coding regions, non-coding regions, or both. Especially preferred are polynucleotide variants containing alterations which produce silent substitutions, additions, or deletions, but do not alter the properties or activities of the encoded polypeptide. Nucleotide variants produced by silent substitutions due to the degeneracy of the genetic code are preferred. CLASP-2 polynucleotide variants can be produced for a variety of reasons, e.g., to optimize codon expression for a particular host (change codons in the human mRNA to those preferred by a bacterial host such as E. coli).

Exemplary CLASP-2 polynucleotide fragments are preferably at least about 15 nucleotides, and more preferably at least about 20 nucleotides, still more preferably at least about 30 nucleotides, and even more preferably, at least about 40 nucleotides in length, or larger 50, 150, 200, 250, 300, 350, 400, 450, 500, 550, 600, 650 nucleotides. In one embodiment, exemplary fragments include fragments having at least a sequence from about nucleotide number 1-50, 51-100, 101-150, 151-200, 201-250, 251-300, 301-350, 351-400, 401-450, 451-500, 501-550, 551-600 to the end of SEQ ID NO: 1 or SEQ ID NO: \_\_\_ or comprising the cDNA coding sequence in the deposited clones. In this context “about” includes the particularly recited ranges, larger or smaller by several (5, 4, 3, 2, or 1) nucleotides, at either terminus or at both termini. Preferably, these fragments encode a polypeptide which has biological activity. More preferably, these polynucleotides can be used as probes or primers as discussed herein.

In other embodiments, CLASP-2 polynucleotides of the invention are other than SEQ ID NO:1 or fragments of SEQ ID NO:1.

As shown in FIG 11 above, there are at least three CLASP-2 full length cDNA isoforms (A + Z, B + Z, and C + Z). Each of the isoforms uses a unique first exon (labelled exon 1A, 1B, and 1C) (see FIG. 11 and Table 4 below).

Table 4: CLASP-2 Isoforms

CLASP-2 Isoform	FIG 11C Schematic	Nucleotides
Isoform 1	A + Z	−182 to 6690
Isoform 2	B + Z	−219 to 6690
Isoform 3	C + Z	−143 to 6690

In one embodiment, the CLASP-2 polynucleotide has the sequence shown in FIG. 11 (Isoform 1, Isoform 2, or Isoform 3 as indicated in Table 4 above) or a fragment of the sequence shown in FIG. 11 comprising at least about 1, 5, 10, 25 or 50 or more contiguous nucleotides from nucleotides −182 to 1883 of Isoform 1, nucleotides −219 to 1883 of Isoform 2, or nucleotides −143 to 1883 of Isoform 3.

In another embodiment, CLASP-2 primers or probes comprise at least about 5, 10, 25 or 50 or more contiguous nucleotides from nucleotides −182 to 1883 of Isoform 1, nucleotides −219 to 1883 of Isoform 2, or nucleotides −143 to 1883 of Isoform 3 as shown in FIG. 11 and Table 4 above alone or in combination with SEQ ID NO:1 or a fragment of SEQ ID NO:1.

In an aspect, the invention provides antibodies or binding fragments that bind the CLASP-2 isoforms 1-3. In another embodiment, the invention provides antibodies that specifically bind to the CLASP-2 isoforms shown in FIG. 11 but not to the polypeptide encoded by SEQ ID NO:1

In one embodiment, the CLASP-2 variants differ from those shown in FIG. 1 or FIG. 11 (SEQ ID NOs 1, 3, 5, 7 9, \_\_\_\_\_) by virtue of incorporating a different combination of exons than found in the exemplified sequences. For example, 81g01 (Genbank Accession Number AF85864; Locus HUMYN81g01; 526 bp; EST sequence submitted August 29, 1998 by Washington University at St. Louis; see FIG. 3A and FIG. 3B) is a variant of hCLASP-2 on the basis of CLASP-2 identity along certain stretches of the sequence. From 5' to 3', it begins with a 315 nucleotide stretch which is identical to CLASP-2A. It then has a gap relative to CLASP-2A that is identical to the GAP in another CLASP-2 isoform, hCLASP-2D (KIAA1058). In place of that gap, a 16 amino acid insert (48 nucleotides) is present which is not found in other isoforms, followed by another

approximately 150 bp stretch of nucleotides once again identical to CLASP-2A. This is characteristic of an alternate splice due to the specific sequence identity on both sides of a differential stretch of nucleotides.

Using known methods of protein engineering and recombinant DNA technology, variants can be generated to improve or alter the characteristics of the CLASP-2 polypeptides. For instance, one or more amino acids can be deleted from the N-terminus or C-terminus of the CLASP-2 protein without substantial loss of biological function.

Furthermore, even if deleting one or more amino acids from the N-terminus or C-terminus of a polypeptide results in modification or loss of one or more biological functions, other biological activities can still be retained. For example, the ability of a deletion variant to induce and/or to bind antibodies which recognize the secreted form will likely be retained when less than the majority of the residues of the secreted form are removed from the N-terminus or C-terminus. Whether a particular polypeptide lacking N or C-terminal residues of a protein retains such immunogenic activities can readily be determined by routine methods described herein and otherwise known in the art.

Thus, the invention further includes CLASP-2 polypeptide variants which show biological activity. Such variants include deletions, insertions, inversions, repeats, and substitutions selected according to general rules known in the art so as to have little effect on activity. For example, guidance concerning how to make phenotypically silent amino acid substitutions is provided in Bowie, J. U. *et al.*, Science 247: 1306-1310 (1990), wherein the authors indicate that there are two main strategies for studying the tolerance of an amino acid sequence to change.

The first strategy exploits the tolerance of amino acid substitutions by natural selection during the process of evolution. By comparing amino acid sequences in different species, conserved amino acids can be identified. These conserved amino acids are likely important for protein function. In contrast, the amino acid positions where substitutions have been tolerated by natural selection indicate that these positions are not critical for protein function. Thus, positions tolerating amino acid substitution could be modified while still maintaining biological activity of the protein.

The second strategy uses genetic engineering to introduce amino acid changes at 30 specific positions of a cloned gene to identify regions critical for protein function. For example, site directed mutagenesis or alanine-scanning mutagenesis (introduction of single alanine mutations at every residue in the molecule) can be used. (Cunningham and Wells,

1989, Science 244: 1081-1085) The resulting mutant molecules can then be tested for biological activity.

In various embodiments, CLASP-2 polynucleotide fragments include coding regions for, or regions hybridizable to, the CLASP-2 structural or functional domains described *supra*. As set out in the Figures, such preferred regions include the following domains/motifs: ITAM, DOCK, COILED/COILED, and PBM. Thus, for example, polypeptide fragments of CLASP-2 as shown in FIG. 1 and FIG. 11-(SEQ ID NO: 2, 4, 6 10, \_\_\_\_\_) falling within conserved domains are specifically contemplated by the present invention (see FIG. 3). Moreover, polynucleotide fragments encoding these domains are also contemplated. Such polypeptide fragments find use, for example, as inhibitors of CLASP-2 function in CLASP-2-expressing cells.

#### **5.4.2. Uses of CLASP-2 Polynucleotides**

The CLASP-2 polynucleotides of the invention are useful in a variety of applications. In one aspect of the invention, the polypeptide-encoding CLASP-2 polynucleotides of the invention are used to express CLASP-2 polypeptides (*e.g.*, as described herein) for example to produce anti-CLASP-antibodies or for use as therapeutic polypeptides. In another aspect, the CLASP-2 polynucleotide or fragments thereof can be used for diagnostic purposes (*e.g.*, as probes for CLASP-2 expression). In particular, since CLASP-2s can be expressed in lymphocytes, a CLASP-2 polynucleotide can be used to detect the expression of CLASP-2 as a lymphocyte marker. For diagnostic purposes, a CLASP-2 polynucleotide can be used to detect CLASP-2 gene expression or aberrant CLASP-2 gene expression in disease states. In another aspect, the CLASP-2 polynucleotide or fragments are used for therapeutic purposes. For example, included in the scope of the invention are methods for inhibiting CLASP-2 expression, *e.g.*, using oligonucleotide sequences, such as antisense RNA and DNA molecules and ribozymes, that function to inhibit expression of CLASP-2. In another aspect, CLASP-2 polynucleotides can be used to construct transgenic and knockout animals, *e.g.*, for screening of CLASP-2 agonists and antagonists. In another aspect, CLASP-2 polynucleotides can be used for screening of CLASP-2 agonists and antagonists.

##### **5.4.2.1. Use of CLASP-2 Polynucleotides for Detection, Diagnosis, and Treatment**

The CLASP-2 polynucleotides of the invention are useful for detection of CLASP-2 expression in cells and in the diagnosis of diseases or disorders (*e.g.*,



immunodeficient states) resulting from aberrant expression of CLASP-2. Aberrant expression of CLASP-2 mRNA or protein means expression in lymphocytes (*e.g.*, T lymphocytes or B lymphocytes) or other CLASP-2 expressing cells of at least 2-fold, preferably at least 5-fold greater or less than expression in control lymphocytes obtained from a healthy subject. CLASP-2 polypeptide expression is easily measured by ELISA using anti-CLASP-2 antibodies of the invention. CLASP-2 mRNA expression (including expression of specific species or splice variants of CLASP-2) can be measured by quantitative Northern analysis or quantitative PCR, LCR, or other methods, using the probes and primers of the invention.

In one embodiment, the assays of the present invention are amplification-based assays for detection of an CLASP-2 gene product. In an amplification based assay, all or part of a CLASP-2 mRNA or cDNA (hereinafter also referred to as "target") is amplified, and the amplification product is then detected directly or indirectly. When there is no underlying gene product to act as a template, no amplification product is produced (*e.g.*, of the expected size), or amplification is non-specific and typically there is no single amplification product. In contrast, when the underlying gene or gene product is present, the target sequence is amplified, providing an indication of the presence and/or quantity of the underlying gene or mRNA. Target amplification-based assays are well known to those of skill in the art.

The present invention provides a wide variety of primers and probes for detecting CLASP-2 genes and gene products. Such primers and probes are sufficiently complementary to the CLASP-2 gene or gene product to hybridize to the target nucleic acid. Primers are typically at least 6 bases in length, usually between about 10 and about 100 bases, typically between about 12 and about 50 bases, and often between about 14 and about 25 bases in length, often PCR primers of 15-30 (*e.g.*, 18-22 nucleotides) are used. However, the length of primers can be adjusted by one skilled in the art. One of skill, having reviewed the present disclosure, will be able, using routine methods, to select primers to amplify all, or any portion, of the CLASP-2 gene or gene product, or to distinguish between variant gene products, CLASP-2 alleles, and the like. Single oligomers (*e.g.*, U.S. Pat. No. 5,545,522), nested sets of oligomers, or even a degenerate pool of oligomers can be employed for amplification.

It will be appreciated that probes and primers can be selected to distinguish between species and splice variants based on the guidance of this disclosure, by targeting

primers or probes to differentially used exons (or exon-exon junctions that differ between variants).

Methods can include the steps of collecting a sample of cells from a patient, isolating nucleic acid (*e.g.*, genomic, mRNA or both) from the cells of the sample, contacting the nucleic acid sample with one or more primers which specifically hybridize to an CLASP-2 gene under conditions such that hybridization and amplification of the CLASP-2-gene (if present) occurs, and detecting the presence or absence of an amplification product, or detecting the size of the amplification product and comparing the length to a control sample. See U.S. Pat. Nos. 4,683,195 and 4,683,202, Landegran *et al.*, 1988, Science 241: 1077-1080; Nakazawa *et al.*, 1994, Proc. Natl. Acad. Sci. U.S.A. 91: 360-364, Abravaya *et al.*, 1995, Nucleic Acids Res. 23: 675-682).

Because CLASP-2 gene products are expressed in the immune system (*e.g.*, T lymphocytes, B lymphocytes and macrophages), expression will be typically assayed in these cells. Methods which are well known to those skilled in the art can be used to isolate lymphocytes, macrophages, and alike (*See, e.g.*, Coligan, J. E., *et al.* (eds.), 1991, Current Protocols in Immunology, John Wiley & Sons, NY; this reference is incorporated by reference for all purposes). In one embodiment, assays are carried out on biopsy or autopsy-derived tissue.

In various embodiments, CLASP-2 gene expression is detected by hybridization of a detectable probe to mRNA or cDNA obtained from cells (*e.g.*, lymphocytes). A variety of methods for specific DNA and RNA measurement using nucleic acid hybridization techniques are known to those of skill in the art (see Sambrook *et al.*, *supra*). Hybridization based assays refer to assays in which a probe nucleic acid is hybridized to a target nucleic acid, forming a hybridization complex. Usually the nucleic acid hybridization probes of the invention are entirely or substantially identical to a contiguous sequence of the CLASP-2 gene or RNA sequence. Preferably, nucleic acid probes are at least about 50 bases, often at least about 20 bases, and sometimes at least about 200 bases, at least about 300-500 nucleotides or more in length. Various hybridization techniques are well known in the art, and are in fact the basis of many commercially available diagnostic kits.

Methods of selecting nucleic acid probe sequences for use in nucleic acid hybridization are discussed in Sambrook *et al.*, *supra*. In some formats, at least one of the target and probe is immobilized. The immobilized nucleic acid can be DNA, RNA, or another oligo- or poly-nucleotide, and can comprise natural or non-naturally occurring

nucleotides, nucleotide analogs, or backbones. Such assays can be in any of several formats including: Southern, Northern, dot and slot blots, high-density polynucleotide or oligonucleotide arrays (e.g., GeneChips<sup>TM</sup> Affymetrix), dip sticks, pins, chips, or beads. All of these techniques are well known in the art and are the basis of many commercially available diagnostic kits. Hybridization techniques are generally described in Hames *et al.*, ed., 1985, Nucleic Acid Hybridization, A Practical Approach IRL Press; Gall and Pardue, 1969, Proc. Natl. Acad. Sci. U.S.A., 63: 378-383; and John *et al.*, 1969, Nature, 223: 582-587.

A variety of nucleic acid hybridization formats are known to those skilled in the art. For example, one common format is direct hybridization, in which a target nucleic acid is hybridized to a labeled, complementary probe. Typically, labeled nucleic acids are used for hybridization, with the label providing the detectable signal. One method for evaluating the presence, absence, or quantity of CLASP-2 mRNA is carrying out a Northern transfer of RNA from a sample and hybridization of a labeled CLASP-2 specific nucleic acid probe. A useful method for evaluating the presence, absence, or quantity of DNA encoding CLASP-2 proteins in a sample involves a Southern transfer of DNA from a sample and hybridization of a labeled CLASP-2 specific nucleic acid probe.

Other common hybridization formats include sandwich assays and competition or displacement assays. Sandwich assays are commercially useful hybridization assays for detecting or isolating nucleic acid sequences. Such assays utilize a “capture” nucleic acid covalently immobilized to a solid support and a labeled “signal” nucleic acid in solution. The biological or clinical sample will provide the target nucleic acid. The “capture” nucleic acid and “signal” nucleic acid probe hybridize with the target nucleic acid to form a “sandwich” hybridization complex. To be effective, the signal nucleic acid cannot hybridize with the capture nucleic acid.

In one embodiment, CLASP-2 polypeptides or polynucleotides are useful in treating deficiencies or disorders of the immune system, by activating or inhibiting the activation, differentiation of immune cells. Immune cells develop through a process called hematopoiesis, producing myeloid (platelets, red blood cells, neutrophils, and macrophages) and lymphoid (B and T lymphocytes) cells from pluripotent stem cells. The etiology of these immune deficiencies or disorders can be genetic, somatic, such as cancer or some autoimmune disorders, acquired (e.g., by chemotherapy or toxins), or infectious.

In another embodiment, CLASP-2 polynucleotides or polypeptides are useful in treating or detecting deficiencies or disorders of hematopoietic cells. CLASP-2

polypeptides or polynucleotides could be used to increase differentiation and proliferation of hematopoietic cells, including the pluripotent stem cells, in an effort to treat those disorders associated with a decrease in certain (or many) types hematopoietic cells. Examples of immunologic deficiency syndromes include, but are not limited to: blood protein disorders (e.g., agammaglobulinemia, dysgammaglobulinemia), ataxia telangiectasia, common variable immunodeficiency, Digeorge Syndrome, HIV infection, HTLV-BLV infection, leukocyte adhesion deficiency syndrome, lymphopenia, phagocyte bactericidal dysfunction, severe combined immunodeficiency (SCIDs), Wiskott-Aldrich Disorder, anemia, thrombocytopenia, or hemoglobinuria.

In one embodiment, CLASP-2 polynucleotides or polypeptides are useful in treating or detecting autoimmune diseases. The term "autoimmune disease" as used herein has the normal meaning in the art and refers to a spontaneous or induced malfunction of the immune system of mammals in which the immune system fails to distinguish between foreign immunogenic substances within the mammal and/or autologous ("self") substances and, as a result, treats autologous ("self") tissues and substances as if they were foreign and mounts an immune response against them. Autoimmune disease is characterized by production of either antibodies that react with self tissue, and/or the activation of immune effector T cells that are autoreactive to endogenous self antigens. Three main immunopathologic mechanisms act to mediate autoimmune diseases: 1) autoantibodies are directed against functional cellular receptors or other cell surface molecules, and either stimulate or inhibit specialized cellular function with or without destruction of cells or tissues; 2) autoantigen--autoantibody immune complexes form in intercellular fluids or in the general circulation and ultimately mediate tissue damage; and 3) lymphocytes produce tissue lesions by release of cytokines or by attracting other destructive inflammatory cell types to the lesions. These inflammatory cells in turn lead to production of lipid mediators and cytokines with associated inflammatory disease.

Since many autoimmune disorders result from inappropriate recognition of self as foreign material by immune cells. This inappropriate recognition results in an immune response leading to the destruction of the host tissue. Therefore, the administration of CLASP-2 polypeptides or polynucleotides that can inhibit an immune response, particularly the proliferation, or differentiation of T-cells, can be an effective therapy in preventing autoimmune disorders.

Examples of autoimmune disorders that can be treated or detected by CLASP-2 include, but are not limited to: Addison's Disease, hemolytic anemia, antiphospholipid

syndrome, rheumatoid arthritis, dermatitis, allergic encephalomyelitis, glomerulonephritis, Goodpasture's Syndrome, Graves' Disease, Multiple Sclerosis, Myasthenia Gravis, Neuritis, Ophthalmia, Bullous Pemphigoid, Pemphigus, Polyendocrinopathies, Purpura, Reiter's Disease, Stiff-Man Syndrome, Autoimmune Thyroiditis, Systemic Lupus Erythematosus, Autoimmune Pulmonary Inflammation, Guillain-Barre Syndrome, insulin dependent diabetes mellitus, and autoimmune inflammatory eye disease.

Similarly, allergic reactions and conditions, such as asthma (particularly allergic asthma) or other respiratory problems, can also be treated by CLASP-2 polypeptides or polynucleotides. Moreover, CLASP-2 can be used to treat anaphylaxis or hypersensitivity to an antigenic molecules.

In one embodiment CLASP-2 polynucleotides or polypeptides are used to treat and/or prevent organ rejection or graft-versus-host disease (GVHD). Organ rejection occurs by host immune cell destruction of the transplanted tissue through an immune response. Similarly, an immune response is also involved in GVHD, but, in this case, the foreign transplanted immune cells destroy the host tissues. The administration of CLASP-2 polypeptides or polynucleotides that inhibits an immune response, particularly the proliferation, differentiation of T-cells, can be an effective therapy in preventing organ rejection or GVHD.

Similarly, in another embodiment, CLASP-2 polypeptides or polynucleotides are used to modulate inflammation. The term "inflammation" refers to both acute responses (*i.e.*, responses in which the inflammatory processes are active) and chronic responses (*i.e.*, responses marked by slow progression and formation of new connective tissue). Acute and chronic inflammation can be distinguished by the cell types involved. Acute inflammation often involves polymorphonuclear neutrophils; whereas chronic inflammation is normally characterized by a lymphohistiocytic and/or granulomatous response. Inflammation includes reactions of both the specific and non-specific defense systems. A specific defense system reaction is a specific immune system reaction response to an antigen (possibly including an autoantigen). A non-specific defense system reaction is an inflammatory response mediated by leukocytes incapable of immunological memory. Such cells include granulocytes, macrophages, neutrophils and eosinophils.

For example, CLASP-2 polypeptides or polynucleotides can inhibit the proliferation and differentiation of cells involved in an inflammatory response. These molecules can be used to treat inflammatory conditions, both chronic and acute conditions, including inflammation associated with infection (*e.g.*, septic shock, sepsis, or systemic



inflammatory response syndrome (SIRS)), ischemia-reperfusion injury, endotoxin lethality, arthritis, complement-mediated hyperacute rejection, nephritis, cytokine or chemokine induced lung injury, inflammatory bowel disease, Crohn's disease, or resulting from over production of cytokines (e.g., TNF or IL-1.). Examples of specific types of inflammation are

5 diffuse inflammation, focal inflammation, croupous inflammation, interstitial inflammation, obliterative inflammation, parenchymatous inflammation, reactive inflammation, specific inflammation, toxic inflammation and traumatic inflammation.

In another embodiment CLASP-2 polypeptides or polynucleotides are used to treat or detect infectious agents. For example, by increasing the immune response,

10 particularly increasing the proliferation and differentiation of B and/or T cells, infectious diseases can be treated. The immune response can be increased by either enhancing an existing immune response, or by initiating a new immune response. CLASP-2 polypeptides or polynucleotides can be used to treat or detect any of these symptoms or diseases.

#### **5.4.2.2. Use of CLASP-2 Polynucleotides in Screening**

The presence or absence of hCLASP-2 nucleotide and amino acid sequences

15 in a biological sample can be used in screening assays as medical diagnostics to aid in clinical decision-making. In one embodiment, hCLASP-2-based diagnostics involves screening assays for vaginal bleeding of unknown cause. In several examples discussed below, the cause of the bleeding can be in part differentiated by knowledge of whether the vaginal

20 bleeding contains placental components (Hart FD, Ed., 1985, French's Index of Differential Diagnosis, 12th Ed. John Wright & Sons, pp. 561-63). In these cases, the high expression of hCLASP-2 nucleotide sequences in placenta relative to its low expression in blood (FIG. 4A) will allow the detection of the presence of placenta based on the presence of the hCLASP-2 nucleotide or protein. Such detection can be achieved by quantitative RT-PCR, Northern

25 analysis, Western analysis, ELISAs, and fluorescence activated cell sorting (FACS) by using labeled anti-hCLASP-2 antibodies (Sambrook *et al.*, 1989, Molecular Cloning, 2nd Ed., Cold Spring Harbor Lab. Press; Harlow *et. al.*, 1988, Antibodies, a laboratory manual, Cold Spring Harbor Lab. Press).

For example, hCLASP-2 can be used in the following screening assays:

30 (1) A woman gives birth and presents with post-partum bleeding. In this case the presence of placental tissue indicates a condition called "retained products of

conception” that requires surgical evacuation of the uterus (Decherney and Pernol, Eds., 1996, Current Obstetric & Gynecologic Diagnosis & Treatment, 8th Ed. McGraw Hill).

(2) A pregnant woman suffers from vaginal bleeding of unknown origin. In this case the presence of placental tissue indicates a condition called “threatened abortion” that implies a poor prognosis for carrying the fetus to term (Decherney and Pernol, Eds., 1996, Current Obstetric & Gynecologic Diagnosis & Treatment, 8th Ed. McGraw Hill).

(3) A woman of child bearing age presents with vaginal bleeding and is found to have a positive pregnancy test without evidence of an intra-uterine pregnancy. In this case, the most serious of the differential diagnoses is ectopic pregnancy, a medical emergency. However, another common diagnosis is a completed abortion or miscarriage. The presence of products of conception (*i.e.* placenta) in the vaginal bleeding strongly favors the diagnosis of completed abortion over that of ectopic pregnancy (Decherney and Pernol, Eds., 1996, Current Obstetric & Gynecologic Diagnosis & Treatment, 8th Ed. McGraw Hill).

In another embodiment, hCLASP-2-based diagnostics involve screening assays to determine injury to vital tissues that express hCLASP-2 at high levels. Such tissues include kidney, heart, and lung (Fig 4A). Injury to these tissues can result in leakage of cells and cellular constituents including hCLASP-2 into surrounding fluids (specified below). Detection of abnormally high levels of hCLASP-2 protein in these surrounding fluids by Western analysis or ELISA, or detection of abnormally high levels of hCLASP-2 RNA in these fluids by RT-PCR or Northern analysis is expected to aid in the diagnosis of tissue injury.

In the case of renal injury, the hCLASP-2 nucleotide or amino acid sequences or fragments thereof would be expected to appear in the urine. Detection of abnormally high levels of hCLASP-2 can aid in the diagnosis of both nephritis and tubular necrosis, and differentiate from non-renal causes of proteinuria. Early diagnosis of nephritis is of particular value in patients with clinical signs and symptoms suggestive of systemic lupus erythematosus in whom early diagnosis and treatment of lupus nephritis can prevent irreversible kidney damage (Cameron J.S., 1999, J Nephrol 12 Suppl 2: S29-41). While tubular necrosis currently cannot be reversed by pharmacotherapy, differentiation of tubular necrosis from pre-renal failure is critical in formulating a treatment plan for oligouric hospitalized patients (Bidani A. and Churchill P.C., 1989, Dis Mon 35: 57-132).

In the case of myocardial injury, the hCLASP-2 nucleic or amino acid sequence or fragments thereof are expected to appear in the blood. This is analogous to current standard practice of monitoring for other elevated levels myocardial proteins (*e.g.*,

creatine kinase, troponin) in the blood following myocardial infarction and ischemia by standard ELISA or electrophoretic methodologies (Fauci *et al.*, (eds.), 1998, Harrison's Principles of Internal Medicine, 14th Ed., McGraw Hill, pp. 1352-1375). The presence of hCLASP-2 in cardiac muscle and its absence in skeletal muscle and blood makes hCLASP-2 an ideal marker to diagnose and monitor myocardial injury.

Unlike myocardial injury, pulmonary injury is not routinely diagnosed by assaying serum for lung-specific proteins. By analogy to myocardial infarction, pulmonary infarction also releases lung-specific proteins and cells into systemic circulation. Pulmonary infarction due to pulmonary embolism (PE) or pneumonia is expected to release hCLASP-2-bearing cells or protein/peptides into systemic circulation. Detection of hCLASP-2 protein in serum or RNA in blood can aid in the diagnosis of pulmonary infarction in the appropriate clinical setting. Current methods to diagnose PE are not only expensive but lack specificity and sensitivity, and the misdiagnosis of this condition is a leading cause of preventable death in hospitalized patients (Raskob G.E. and Hull R.D., 1999, Curr Opin Hematol. 6(5): 280-4).

In another embodiment, hCLASP-2-based diagnostics involve screening assays for identifying disorders of cells of hematopoietic lineage. hCLASP-2 is expressed in human T cells, B cells but not cells from the myeloid lineage. Different hCLASP-2 isoforms in T and B cells permit further discrimination between malignancies of T and B lineage (FIG. 4B). Precise identification of hematopoietic cell types is vital to guide chemotherapy and radiation therapy of patients with leukemia and lymphoma (Fauci et al Eds., 1998, Harrison's Principles of Internal Medicine, 14th Ed. McGraw Hill, pp. 695-712). hCLASP-2 expression differences can be detected by using FACS, immunofluorescence, immunoperoxidase staining, RT-PCR, in situ hybridization or RNA blot analysis (Sambrook, Fritsch and Maniatis, Molecular Cloning, 2nd Ed. Cold Spring Harbor Lab. Press, 1989; Ward MS, Pathology 1999 Nov; 31(4): 382-92).

In another embodiment, hCLASP-2-based diagnostics involve screening assays for identifying activated immune system cells. Although hCLASP-2 is generally expressed at quite low levels in PBMCs (which is critical for some of the above applications), it is known that the surface expression of the closely related mouse CLASP-1 protein is altered during the process of lymphocyte activation. An analogous change in expression is expected for the hCLASP-2 protein. Subtyping lymphocytes specific for a particular antigen, for example, using MHC-based multimeric staining reagents (Altman et. al., 1996, Science 274: 94-6), for separating cell populations into hCLASP-2 high and hCLASP-2 low populations, can aid in determining the nature of the immune response against that antigen.

Such understanding is critical, for example, in predicting the course of chronic viral infections such as hepatitis B, hepatitis C, and HIV, and to designing appropriate treatment regimens for patients suffering from these infections.

hCLASP-2 can also serve as a potential therapeutic agent for Wilms' tumor.

5 Wilms tumor is the most common primary renal tumor of childhood (Cotran, Kumar, and Collins, 1999, Robbins Pathologic Basis of Disease, 6th Ed. W.B. Saunders, pp. 487-89). As discussed herein, hCLASP-2 is highly expressed in 293 cells, embryonic kidney epithelial cells. Therefore, hCLASP-2 nucleic or amino acid sequence or fragments can serve as tumor markers for Wilms' tumor. Antibodies directed against a hCLASP-2 variant that is expressed  
10 only in Wilms' tumor can serve as novel therapeutic agents for Wilms' tumor, and can also function as delivery vehicles for other targeted therapeutics that may be attached to the anti-hCLASP-2 antibody (*e.g.*, chemotherapeutics or radiolabeling).

#### **5.4.2.2.1. CLASP-2 Antisense, Ribozyme and Triplex Polynucleotides and Methods of Use**

15 Oligonucleotide sequences, that include anti-sense RNA and DNA molecules and ribozymes that function to inhibit the translation of a CLASP-2 mRNA are within the scope of the invention. Such molecules are useful in cases where downregulation of CLASP-2 expression is desired. Anti-sense RNA and DNA molecules act to directly block the translation of mRNA by binding to targeted mRNA and preventing protein translation. The  
20 invention provides methods and antisense oligonucleotide or polynucleotide reagents which can be used to reduce expression of CLASP-2 gene products in vitro or in vivo.

Administration of the antisense reagents of the invention to a target cell results in reduced CLASP activity. As will be apparent to one of skill and as discussed *supra* (Table 3), specific CLASP-2 splice variants can be specifically targeted for inhibition. Alternatively, by  
25 designing an, *e.g.*, antisense molecule that recognizes a sequence found in several or all CLASP-2 species, a general inhibition can be achieved.

#### **A. Antisense**

Without intending to be limited to any particular mechanism, it is believed that antisense oligonucleotides bind to, and interfere with the translation of, the sense CLASP-2  
30 mRNA. Alternatively, the antisense molecule can render the CLASP-2 mRNA susceptible to nuclease digestion, interfere with transcription, interfere with processing, localization or otherwise with RNA precursors ("pre-mRNA"), repress transcription of mRNA from the

CLASP-2 gene, or act through some other mechanism. However, the particular mechanism by which the antisense molecule reduces CLASP-2 expression is not critical.

The antisense polynucleotides of the invention comprise an antisense sequence of at least 7 to 10 to typically 20 or more nucleotides that specifically hybridize to a sequence from mRNA encoding CLASP-2 or mRNA transcribed from the CLASP-2 gene. More often, the antisense polynucleotide of the invention is from about 10 to about 50 nucleotides in length or from about 14 to about 35 nucleotides in length. In other embodiments, antisense polynucleotides are polynucleotides of less than about 100 nucleotides or less than about 200 nucleotides. In general, the antisense polynucleotide should be long enough to form a stable duplex but short enough, depending on the mode of delivery, to administer in vivo, if desired. The minimum length of a polynucleotide required for specific hybridization to a target sequence depends on several factors, such as G/C content, positioning of mismatched bases (if any), degree of uniqueness of the sequence as compared to the population of target polynucleotides, and chemical nature of the polynucleotide (*e.g.*, methylphosphonate backbone, peptide nucleic acid, phosphorothioate), among other factors. Generally, to assure specific hybridization, the antisense sequence is substantially complementary to the target CLASP-2 mRNA sequence. In certain embodiments, the antisense sequence is exactly complementary to the target sequence. The antisense polynucleotides can also include, however, nucleotide substitutions, additions, deletions, transitions, transpositions, or modifications, or other nucleic acid sequences or non-nucleic acid moieties so long as specific binding to the relevant target sequence corresponding to CLASP-2 RNA or its gene is retained as a functional property of the polynucleotide.

It will be appreciated that the CLASP-2 polynucleotides and oligonucleotides of the invention can be made using nonstandard bases (*e.g.*, other than adenine, cytidine, guanine, thymine, and uridine) or nonstandard backbone structures to provides desirable properties (*e.g.*, increased nuclease-resistance, tighter-binding, stability or a desired TM). Techniques for rendering oligonucleotides nuclease-resistant include those described in PCT publication WO 94/12633. A wide variety of useful modified oligonucleotides may be produced, including oligonucleotides having a peptide-nucleic acid (PNA) backbone (Nielsen *et al.*, 1991, Science 254: 1497) or incorporating 2'-O-methyl ribonucleotides, phosphorothioate nucleotides, methyl phosphonate nucleotides, phosphotriester nucleotides, phosphorothioate nucleotides, phosphoramidates. Still other useful oligonucleotides may contain alkyl and halogen-substituted sugar moieties comprising one of the following at the 2' position: OH, SH, SCH<sub>3</sub>, F, OCN, OCH<sub>3</sub>OCH<sub>3</sub>, OCH<sub>3</sub>O(CH<sub>2</sub>)<sub>n</sub>CH<sub>3</sub>, O(CH<sub>2</sub>)<sub>n</sub>NH<sub>2</sub> or



O(CH<sub>2</sub>)<sub>n</sub>CH<sub>3</sub>, where n is from 1 to about 10; C<sub>1</sub> to C<sub>10</sub> lower alkyl, substituted lower alkyl, alkaryl or aralkyl; Cl; Br; CN; CF<sub>3</sub>; OCF<sub>3</sub>; O-, S-, or N-alkyl; O-, S-, or N-alkenyl; SOCH<sub>3</sub>; SO<sub>2</sub>CH<sub>3</sub>; ONO<sub>2</sub>; NO<sub>2</sub>; N<sub>3</sub>; NH<sub>2</sub>; heterocycloalkyl; heterocycloalkaryl; aminoalkylamino; polyalkylamino; substituted silyl; an RNA cleaving group; a cholesteryl group; a folate group; a reporter group; an intercalator; a group for improving the pharmacokinetic properties of an oligonucleotide; or a group for improving the pharmacodynamic properties of an oligonucleotide and other substituents having similar properties. Folate, cholesterol or other groups that facilitate oligonucleotide uptake, such as lipid analogs, may be conjugated directly or via a linker at the 2' position of any nucleoside or at the 3' or 5' position of the 3'-terminal or 5'-terminal nucleoside, respectively. One or more such conjugates may be used. Oligonucleotides may also have sugar mimetics such as cyclobutyls in place of the pentofuranosyl group. Other embodiments may include at least one modified base form or "universal base" such as inosine, or inclusion of other nonstandard bases such as queosine and wybutosine as well as acetyl-, methyl-, thio- and similarly modified forms of adenine, cytidine, guanine, thymine, and uridine which are not as easily recognized by endogenous endonucleases. The antisense oligonucleotide can comprise at least one modified base moiety which is selected from the group including, but not limited to, 5-fluorouracil, 5-bromouracil, 5-chlorouracil, 5-iodouracil, hypoxanthine, xanthine, 4-acetylcytosine, 5-(carboxyhydroxymethyl) uracil, 5-carboxymethylaminomethyl-2-thiouridine, 5-carboxymethylaminomethyluracil, dihydrouracil, beta-D-galactosylqueosine, inosine, N6-isopentenyladenine, 1-methylguanine, 1-methylinosine, 2,2-dimethylguanine, 2-methyladenine, 2-methylguanine, 3-methylcytosine, 5-methylcytosine, N6-adenine, 7-methylguanine, 5-methylaminomethyluracil, 5-methoxyaminomethyl-2-thiouracil, beta-D-mannosylqueosine, 5'-methoxycarboxymethyluracil, 5-methoxyuracil, 2-methylthio-N6-isopentenyladenine, uracil-5-oxyacetic acid (v), wybutoxosine, pseudouracil, queosine, 2-thiocytosine, 5-methyl-2-thiouracil, 2-thiouracil, 4-thiouracil, 5-methyluracil, uracil-5-oxyacetic acid methylester, uracil-5-oxyacetic acid (v), 5-methyl-2-thiouracil, 3-(3-amino-3-N-2-carboxypropyl) uracil, (acp3)w, and 2,6-diaminopurine.

The invention further provides oligonucleotides having backbone analogues such as phosphodiester, phosphorothioate, phosphorodithioate, methylphosphonate, phosphoramidate, alkyl phosphotriester, sulfamate, 3'-thioacetal, methylene(methylimino), 3'-N-carbamate, morpholino carbamate, chiral-methyl phosphonates, nucleotides with short chain alkyl or cycloalkyl intersugar linkages, short chain heteroatomic or heterocyclic

intersugar ("backbone") linkages, or CH<sub>2</sub>-NH-O-CH<sub>2</sub>, CH<sub>2</sub>-N(CH<sub>3</sub>)-OCH<sub>2</sub>, CH<sub>2</sub>-O-N(CH<sub>3</sub>)-CH<sub>2</sub>, CH<sub>2</sub>-N(CH<sub>3</sub>)-N(CH<sub>3</sub>)-CH<sub>2</sub> and O-N(CH<sub>3</sub>)-CH<sub>2</sub>-CH<sub>2</sub> backbones (where phosphodiester is O-P-O-CH<sub>2</sub>), or mixtures of the same. Also useful are oligonucleotides having morpholino backbone structures (U.S. Patent No. 5,034,506).

Useful references include Oligonucleotides and Analogues, A Practical Approach, edited by F. Eckstein, IRL Press at Oxford University Press (1991); Antisense Strategies, Annals of the New York Academy of Sciences, Volume 600, Eds. Baserga and Denhardt (NYAS 1992); Milligan *et al.*, 9 July 1993, J. Med. Chem. 36(14): 1923-1937; Antisense Research and Applications (1993, CRC Press), in its entirety and specifically Chapter 15, by Sanghvi, entitled "Heterocyclic base modifications in nucleic acids and their applications in antisense oligonucleotides;" and Antisense Therapeutics, ed. Sudhir Agrawal (Humana Press, Totowa, New Jersey, 1996).

In one embodiment, the antisense sequence is complementary to relatively accessible sequences of the CLASP-2 mRNA (*e.g.*, relatively devoid of secondary structure). This can be determined by analyzing predicted RNA secondary structures using, for example, the MFOLD program (Genetics Computer Group, Madison WI) and testing in vitro or in vivo as is known in the art. Another useful method for identifying effective antisense compositions uses combinatorial arrays of oligonucleotides (see, *e.g.*, Milner *et al.*, 1997, Nature Biotechnology 15: 537). Examples of oligonucleotides that can be tested in cells for antisense suppression of CLASP-2 function are those capable of hybridizing to (*i.e.*, substantially complementary to) the following positions from SEQUENCE ID NO: 1:

- 1) GAAGGCGATCATCACGTGGCCTTCCATCGC
- 2) GCTTCAAGTAATGACTGGTGCAGAACATCTG
- 3) GCTCCTCCTCAGGCAGGCGCTATGGCTGTGG
- 4) GTAGGCCCGGTGCAGCGTGTGCATACAGATGG

(See also Example 8)

In some embodiments, administration of antisense oligonucleotides will result in reduction of hCLASP-mRNA expression by at least about 50%, as assessed by Northern analysis after administration of an antisense phosphorothioate oligonucleotide at a concentration of 1  $\mu$ M, 5  $\mu$ M, 10  $\mu$ M or 20  $\mu$ M.

The invention also provides an antisense polynucleotide that has sequences in addition to the antisense sequence (*i.e.*, in addition to anti-CLASP-2-sense sequence). In this

case, the antisense sequence is contained within a polynucleotide of longer sequence. In another embodiment, the sequence of the polynucleotide consists essentially of, or is, the antisense sequence.

The antisense nucleic acids (DNA, RNA, modified, analogues, and the like) can be made using any suitable method for producing a nucleic acid, such as the chemical synthesis and recombinant methods disclosed herein. In one embodiment, for example, antisense RNA molecules of the invention can be prepared by *de novo* chemical synthesis or by cloning. For example, an antisense RNA that hybridizes to CLASP-2 mRNA can be made by inserting (ligating) an CLASP-2 DNA sequence (*e.g.*, SEQUENCE ID No: 1, or fragment thereof) in reverse orientation operably linked to a promoter in a vector (*e.g.*, plasmid). Provided that the promoter and, preferably termination and polyadenylation signals, are properly positioned, the strand of the inserted sequence corresponding to the noncoding strand will be transcribed and act as an antisense oligonucleotide of the invention. The term “operably linked” refers to a functional linkage between a nucleic acid expression control sequence (such as a promoter or enhancer) and a second nucleic acid sequence, wherein the expression control sequence directs transcription of the nucleic acid corresponding to the second sequence.

In one embodiment, antisense DNA oligodeoxyribonucleotides derived from the translation initiation site, *e.g.*, between -10 and +10 regions of a CLASP-2 nucleotide sequence, are used. For general methods relating to antisense polynucleotides, see ANTISENSE RNA AND DNA, 1988, D.A. Melton, Ed., Cold Spring Harbor Laboratory, Cold Spring Harbor, NY). See also, Dagle *et al.*, 1991, Nucleic Acids Research, 19: 1805. For a review of antisense therapy, see, *e.g.*, Uhlmann *et al.*, 1990, Chem. Reviews, 90: 543-584.

#### **B. Ribozyme**

Ribozymes are enzymatic RNA molecules capable of catalyzing the specific cleavage of RNA. The mechanism of ribozyme action involves sequence specific hybridization of the ribozyme molecule to complementary target RNA, followed by endonucleolytic cleavage. Within the scope of the invention are engineered hammerhead motif ribozyme molecules that specifically and efficiently catalyze endonucleolytic cleavage of CLASP-2 RNA sequences.

Specific ribozyme cleavage sites within any potential RNA target are initially identified by scanning the target molecule for ribozyme cleavage sites which include the following sequences, GUA, GUU and GUC. Once identified, short RNA sequences of between 15 and 20 ribonucleotides corresponding to the region of the target gene containing

the cleavage site can be evaluated for predicted structural features such as secondary structure that can render the oligonucleotide sequence unsuitable. The suitability of candidate targets can also be evaluated by testing their accessibility to hybridization with complementary oligonucleotides, using ribonuclease protection assays.

### 5 C. Triplex

Alternatively, endogenous target gene expression can be reduced by targeting deoxyribonucleotide sequences complementary to the regulatory region of the target gene (*i.e.*, the target gene promoter and/or enhancers) to form triple helical structures that prevent transcription of the target gene in target cells in the body. (See generally, Helene, 1991, Anticancer Drug Des., 6(6): 569-584; Helene *et al.*, 1992, Ann. N.Y. Acad. Sci., 660: 27-36; and Maher, 1992, Bioassays 14(12): 807-815).

15 Nucleic acid molecules to be used in triplex helix formation for the inhibition of transcription should be single stranded and composed of deoxynucleotides. The base composition of these oligonucleotides must be designed to promote triple helix formation via Hoogsteen base pairing rules, which generally require sizable stretches of either purines or pyrimidines to be present on one strand of a duplex. Nucleotide sequences can be pyrimidine-based, which will result in TAT and CGC<sup>+</sup> triplets across the three associated strands of the resulting triple helix. The pyrimidine-rich molecules provide base complementarily to a purine-rich region of a single strand of the duplex in a parallel orientation to that strand. In addition, nucleic acid molecules can be chosen that are purine-rich, for example, contain a stretch of G residues. These molecules will form a triple helix with a DNA duplex that is rich in GC pairs, in which the majority of the purine residues are located on a single strand of the targeted duplex, resulting in GGC triplets across the three strands in the triplex.

25 Alternatively, the potential sequences that can be targeted for triple helix formation can be increased by creating a so called "switchback" nucleic acid molecule. Switchback molecules are synthesized in an alternating 5'-3', 3'-5' manner, such that they base pair with first one strand of a duplex and then the other, eliminating the necessity for a sizable stretch of either purines or pyrimidines to be present on one strand of a duplex.

### 30 D. General

The anti-sense RNA and DNA molecules, ribozymes and triple helix molecules of the invention can be prepared by any method known in the art for the synthesis of RNA molecules. These include techniques for chemically synthesizing oligodeoxyribonucleotides well known in the art such as for example solid phase

phosphoramidite chemical synthesis. Alternatively, RNA molecules can be generated by *in vitro* and *in vivo* transcription of DNA sequences encoding the antisense RNA molecule. Such DNA sequences can be incorporated into a wide variety of vectors which contain suitable RNA polymerase promoters such as the T7 or SP6 polymerase promoters.

- 5 Alternatively, antisense cDNA constructs that synthesize antisense RNA constitutively or inducibly, depending on the promoter used, can be introduced stably into cell lines.

Various modifications to the DNA molecules can be introduced as a means of increasing intracellular stability and half-life. Possible modifications include, but are not limited to, the addition of flanking sequences of ribo- or deoxy- nucleotides to the 5' and/or  
10 3' ends of the molecule or the use of phosphorothioate or 2' O-methyl rather than phosphodiesterase linkages within the oligodeoxyribonucleotide backbone.

Methods for introducing polynucleotides into such cells or tissue include methods for *in vitro* introduction of polynucleotides such as the insertion of naked polynucleotide, *i.e.*, by injection into tissue, the introduction of a CLASP-2 polynucleotide in  
15 a cell *ex vivo*, the use of a vector such as a virus, (*e.g.*, a retrovirus, adenovirus, adeno-associated virus, and the like), phage or plasmid, and the like or techniques such as electroporation or calcium phosphate precipitation.

#### **5.4.2.2.2. Gene Therapy**

By introducing gene sequences into cells, gene therapy can be used to treat  
20 conditions in which the cells do not express normal CLASP-2 or express abnormal/inactive CLASP-2. In some instances, the polynucleotide encoding a CLASP-2 is intended to replace or act in the place of a functionally deficient endogenous gene. Alternatively, abnormal conditions characterized by overexpression can be treated using the gene therapy techniques described below.

25 In a specific embodiment, nucleic acids comprising a sequence encoding a CLASP-2 protein or functional derivative thereof, are administered to promote CLASP-2 function, by way of gene therapy. Gene therapy refers to therapy performed by the administration of a nucleic acid to a subject. In this embodiment of the invention, the nucleic acid produces its encoded protein that mediates a therapeutic effect by promoting CLASP-2  
30 function.

Any of the methods for gene therapy available in the art can be used according to the present invention. Exemplary methods are described below.



For general reviews of the methods of gene therapy, *see*, Goldspiel *et al.*, 1993, Clinical Pharmacy 12: 488-505; Wu and Wu, 1991, Biotherapy 3: 87-95; Tolstoshev, 1993, Ann. Rev. Pharmacol. Toxicol. 32: 573-596; Mulligan, 1993, Science 260: 926-932; and Morgan and Anderson, 1993, Ann. Rev. Biochem. 62: 191-217; Can, 1993, TIBTECH 11(5): 155-215). Methods commonly known in the art of recombinant DNA technology which can be used are described in Ausubel *et al.*, *supra*; and Kriegler, 1990, Gene Transfer and Expression, A Laboratory Manual, Stockton Press, NY.

In one aspect, the therapeutic composition comprises a CLASP-2 nucleic acid that is part of an expression vector that encodes a CLASP-2 protein or fragment or chimeric protein thereof in a suitable host. In particular, such a nucleic acid has a promoter operably linked to the CLASP-2 coding region, said promoter being inducible or constitutive, and, optionally, tissue-specific. In another particular embodiment, a nucleic acid molecule is used in which the CLASP-2 coding sequences and any other desired sequences are flanked by regions that promote homologous recombination at a desired site in the genome, thus providing for intrachromosomal expression of the CLASP-2 nucleic acid (Koller and Smithies, 1989, Proc. Natl. Acad. Sci. U.S.A. 86: 8932-8935; Zijlstra *et al.*, 1989, Nature 342: 435-438).

Delivery of the nucleic acid into a patient can be either direct, in which case the patient is directly exposed to the nucleic acid or nucleic acid-carrying vector, or indirect, in which case, cells are first transformed with the nucleic acid *in vitro*, then transplanted into the patient. These two approaches are known, respectively, as *in vivo* or *ex vivo* gene therapy.

In a specific embodiment, the nucleic acid is directly administered *in vivo*, where it is expressed to produce the encoded product. This can be accomplished by any of numerous methods known in the art, *e.g.*, by constructing it as part of an appropriate nucleic acid expression vector and administering it so that it becomes intracellular, *e.g.*, by infection using a defective or attenuated retroviral or other viral vector (*see*, U.S. Patent No. 4,980,286), or by direct injection of naked DNA, or by use of microparticle bombardment (*e.g.*, a gene gun; Biolistic, Dupont), or coating with lipids or cell-surface receptors or transfecting agents, encapsulation in liposomes, microparticles, or microcapsules, or by administering it in linkage to a peptide which is known to enter the nucleus, by administering it in linkage to a ligand subject to receptor-mediated endocytosis (*see, e.g.*, Wu and Wu, 1987, J. Biol. Chem. 262: 4429-4432) (which can be used to target cell types specifically expressing the receptors), and the like. In another embodiment, a nucleic acid-ligand

complex can be formed in which the ligand comprises a fusogenic viral peptide to disrupt endosomes, allowing the nucleic acid to avoid lysosomal degradation. In yet another embodiment, the nucleic acid can be targeted *in vivo* for cell specific uptake and expression, by targeting a specific receptor (*see, e.g.*, PCT Publications WO 92/06180 dated April 16, 1992; WO 92/22635 dated December 23, 1992; WO 92/20316 dated November 26, 1992; WO 93/14188 dated July 22, 1993; WO 93/20221 dated October 14, 1993). Alternatively, the nucleic acid can be introduced intracellularly and incorporated within host cell DNA for expression, by homologous recombination (Koller and Smithies, 1989, Proc. Natl. Acad. Sci. U.S.A. 86: 8932-8935; Zijlstra *et al.*, 1989, Nature 342: 435-438).

In a specific embodiment, a viral vector that contains the CLASP-2 nucleic acid is used. For example, a retroviral vector can be used (*see*, Miller *et al.*, 1993, Meth. Enzymol. 217: 581-599). These retroviral vectors have been modified to delete retroviral sequences that are not necessary for packaging of the viral genome and integration into host cell DNA. The CLASP-2 nucleic acid to be used in gene therapy is cloned into the vector, which facilitates delivery of the gene into a patient. More detail about retroviral vectors can be found in Boesen *et al.*, 1994, Biotherapy 6: 291-302, which describes the use of a retroviral vector to deliver the *mdr1* gene to hematopoietic stem cells in order to make the stem cells more resistant to chemotherapy. Other references illustrating the use of retroviral vectors in gene therapy are: Clowes *et al.*, 1994, J. Clin. Invest. 93: 644-651; Kiem *et al.*, 1994, Blood 83: 1467-1473; Salmons and Gunzberg, 1993, Human Gene Therapy 4: 129-141; and Grossman and Wilson, 1993, Curr. Opin. in Genetics and Devel. 3: 110-114.

Adenoviruses are other viral vectors that can be used in gene therapy. Adenoviruses are especially attractive vehicles for delivering genes to respiratory epithelia. Adenoviruses naturally infect respiratory epithelia where they cause a mild disease. Other targets for adenovirus-based delivery systems are liver, the central nervous system, endothelial cells, and muscle. Adenoviruses have the advantage of being capable of infecting non-dividing cells. Kozarsky and Wilson 1993, Current Opinion in Genetics and Development 3: 499-503) present a review of adenovirus-based gene therapy. Bout *et al.*, 1994, Human Gene Therapy 5: 3-10, demonstrated the use of adenovirus vectors to transfer genes to the respiratory epithelia of rhesus monkeys. Other instances of the use of adenoviruses in gene therapy can be found in Rosenfeld *et al.*, 1991, Science 252: 431-434; Rosenfeld *et al.*, 1992, Cell 68: 143-155; and Mastrangeli *et al.*, 1993, J. Clin. Invest. 91: 225-234. Adeno-associated virus (AAV) has also been proposed for use in gene therapy (Walsh *et al.*, 1993, Proc. Soc. Exp. Biol. Med. 204: 289-300).

Another approach to gene therapy involves transferring a gene to cells in tissue culture by such methods as electroporation, lipofection, calcium phosphate mediated transfection, or viral infection. Usually, the method of transfer includes the transfer of a selectable marker to the cells. The cells are then placed under selection to isolate those cells that have taken up and are expressing the transferred gene. Those cells are then delivered to a patient.

In this embodiment, the nucleic acid is introduced into a cell prior to administration *in vivo* of the resulting recombinant cell. Such introduction can be carried out by any method known in the art, including but not limited to transfection, electroporation, microinjection, infection with a viral or bacteriophage vector containing the nucleic acid sequences, cell fusion, chromosome-mediated gene transfer, microcell-mediated gene transfer, spheroplast fusion, and the like. Numerous techniques are known in the art for the introduction of foreign genes into cells (*see, e.g.*, Loeffler and Behr, 1993, Meth. Enzymol. 217: 599-618; Cohen *et al.*, 1993, Meth. Enzymol. 217: 618-644; Cline, 1985, Pharmac. Ther. 29: 69-92) and can be used in accordance with the present invention, provided that the necessary developmental and physiological functions of the recipient cells are not disrupted. The technique should provide for the stable transfer of the nucleic acid to the cell, so that the nucleic acid is expressible by the cell and preferably heritable and expressible by its cell progeny.

The resulting recombinant cells can be delivered to a patient by various methods known in the art. In a preferred embodiment, epithelial cells are injected, *e.g.*, subcutaneously. In another embodiment, recombinant skin cells can be applied as a skin graft onto the patient. Recombinant blood cells (*e.g.*, hematopoietic stem or progenitor cells) are preferably administered intravenously. The amount of cells envisioned for use depends on the desired effect, patient state, and the like., and can be determined by one skilled in the art.

Cells into which a nucleic acid can be introduced for purposes of gene therapy encompass any desired, available cell type, and include but are not limited to epithelial cells, endothelial cells, keratinocytes, fibroblasts, muscle cells, hepatocytes; blood cells such as T lymphocytes, B lymphocytes, monocytes, macrophages, neutrophils, eosinophils, megakaryocytes, granulocytes; various stem or progenitor cells, in particular hematopoietic stem or progenitor cells, *e.g.*, as obtained from bone marrow, umbilical cord blood, peripheral blood, fetal liver, and the like. In a preferred embodiment, the cell used for gene therapy is autologous to the patient.

In a specific embodiment, the nucleic acid to be introduced for purposes of gene therapy comprises an inducible promoter operably linked to the coding region, such that expression of the nucleic acid is controllable by controlling the presence or absence of the appropriate inducer of transcription.

#### 5.4.2.3. Knockout Cells

In one aspect of the invention, endogenous target gene expression can also be reduced by inactivating or “knocking out” the target gene or its promoter using targeted homologous recombination (*see, e.g.*, Smithies *et al.*, 1985, Nature 317: 230-234; Thomas and Capecchi, 1987, Cell 51: 503-512; Thompson *et al.*, 1989, Cell 5: 313-321; each of which is incorporated by reference herein in its entirety). For example, a mutant, non-functional target gene (or a completely unrelated DNA sequence) flanked by DNA homologous to the endogenous target gene (either the coding regions or regulatory regions of the target gene) can be used, with or without a selectable marker and/or a negative selectable marker, to transfect cells that express the target gene *in vivo*. Insertion of the DNA construct, via targeted homologous recombination, results in inactivation of the target gene. Such approaches are particularly suited in the agricultural field where modifications to ES (embryonic stem) cells can be used to generate animal offspring with an inactive target gene (*see, e.g.*, Thomas and Capecchi, 1987 and Thompson, 1989, *supra*). However, this approach can be adapted for use in humans provided the recombinant DNA constructs are directly administered or targeted to the required site *in vivo* using appropriate viral vectors.

#### 5.4.2.4. Transgenic and Knockout Animals

The CLASP-2 gene product can also be expressed in transgenic animals. Animals of any species, including, but not limited to, mice, rats, rabbits, guinea pigs, pigs, micro-pigs, goats, sheep, and non-human primates, *e.g.*, baboons, monkeys, and chimpanzees can be used to generate CLASP-2 transgenic animals. The term “transgenic,” as used herein, refers to animals expressing CLASP-2 gene sequences from a different species (*e.g.*, mice expressing human CLASP-2 gene sequences), as well as animals that have been genetically engineered to overexpress endogenous (*i.e.*, same species) CLASP-2 sequences or animals that have been genetically engineered to no longer express endogenous CLASP-2 gene sequences (*i.e.*, “knock-out” animals), and their progeny.

Any technique known in the art can be used to introduce a CLASP-2 transgene into animals to produce the founder lines of transgenic animals. Such techniques include, but

are not limited to pronuclear microinjection (Hoppe and Wagner, 1989, U.S. Pat. No. 4,873,191); retrovirus mediated gene transfer into germ lines (Van der Putten *et al.*, 1985, Proc. Natl. Acad. Sci., U.S.A. 82: 6148-6152); gene targeting in embryonic stem cells (Thompson *et al.*, 1989, Cell 56: 313-321); electroporation of embryos (Lo, 1983, Mol. Cell. Biol. 3: 1803-1814); and sperm-mediated gene transfer (Lavitrano *et al.*, 1989, Cell 57: 717-723) (For a review of such techniques, see Gordon, 1989, Transgenic Animals, Intl. Rev. Cytol. 115, 171-229)

Any technique known in the art can be used to produce transgenic animal clones containing a CLASP-2 transgene, for example, nuclear transfer into enucleated oocytes of nuclei from cultured embryonic, fetal or adult cells induced to quiescence (Campbell *et al.*, 1996, Nature 380: 64-66; Wilmut *et al.*, Nature 385: 810-813).

The present invention provides for transgenic animals that carry a CLASP-2 transgene in all their cells, as well as animals that carry the transgene in some, but not all their cells, *i.e.*, mosaic animals. The transgene can be integrated as a single transgene or in concatamers, *e.g.*, head-to-head tandems or head-to-tail tandems. The transgene can also be selectively introduced into and activated in a particular cell type by following, for example, the teaching of Lasko *et al.* (1992, Proc. Natl. Acad. Sci. U.S.A. 89: 6232-6236). The regulatory sequences required for such a cell-type specific activation will depend upon the particular cell type of interest, and will be apparent to those of skill in the art. When it is desired that the CLASP-2 transgene be integrated into the chromosomal site of the endogenous CLASP-2 gene, gene targeting is preferred. Briefly, when such a technique is to be utilized, vectors containing some nucleotide sequences homologous to the endogenous CLASP-2 gene are designed for the purpose of integrating, via homologous recombination with chromosomal sequences, into and disrupting the function of the nucleotide sequence of the endogenous CLASP-2 gene. The transgene can also be selectively introduced into a particular cell type, thus inactivating the endogenous CLASP-2 gene in only that cell type, by following, for example, the teaching of Gu *et al.* (1994, Science 265: 103-106). The regulatory sequences required for such a cell-type specific inactivation will depend upon the particular cell type of interest, and will be apparent to those of skill in the art.

Once transgenic animals have been generated, the expression of the recombinant CLASP-2 gene can be assayed utilizing standard techniques. Initial screening can be accomplished by Southern blot analysis or PCR techniques to analyze animal tissues to assay whether integration of the transgene has taken place. The level of mRNA expression of the transgene in the tissues of the transgenic animals can also be assessed using techniques



that include, but are not limited to, Northern blot analysis of tissue samples obtained from the animal, *in situ* hybridization analysis, and RT-PCR (reverse transcriptase PCR). Samples of CLASP-2 gene-expressing tissue, can also be evaluated immunocytochemically using antibodies specific for the CLASP-2 transgene product.

#### 5 **5.4.2.5. Other Uses of CLASP-2 Polynucleotides**

There exists an ongoing need to identify new chromosome marking reagents. Sequences can be mapped to chromosomes by preparing PCR primers from SEQ ID NO: 1, 3, 5, or 9. These primers can be less than 50 nucleotides in length, generally less than 46 nucleotides, more generally less than 41 nucleotides, most generally less than 36 nucleotides, preferably less than 31 nucleotides, more preferably less than 26 nucleotides, and most preferably less than 21 nucleotides in length. The probes can also be less than 16 nucleotides, less than 13 nucleotides in length, less than 9 nucleotides in length and less than 7 nucleotides in length. Primers can be selected so that the primers do not span more than one predicted exon in the genomic DNA. These primers are then used for PCR screening of somatic cell hybrids containing individual human chromosomes (*i.e.*, chromosome 13). Only those hybrids containing the human CLASP-2 gene corresponding to SEQ ID NO: 1, 3, 5, or 9 will yield an amplified fragment.

Similarly, somatic hybrids provide a rapid method of PCR mapping the polynucleotides to particular chromosomes. Precise chromosomal location of the CLASP-2 polynucleotides can also be achieved using fluorescence in situ hybridization (FISH) of a metaphase chromosomal spread. See Verma, et al, Human Chromosomes: A Manual of Basic Techniques, Pergamon Press. NY, 1988. Once a polynucleotide has been mapped to a precise chromosomal location, the physical position of the polynucleotide can be used in linkage analysis. Linkage analysis establishes coinheritance between a chromosomal location and presentation of a particular disease. See McKusick, V., 1998, Mendelian Inheritance in Man : A Catalog of Human Genes and Genetic Disorders, 12th Ed, Johns Hopkins University Press.

The CLASP-2 polynucleotides can be used for identifying individuals from minute biological samples as DNA markers for restriction fragment length polymorphism (RFLP). An individual's genomic DNA is digested with one or more restriction enzymes, and probed on a Southern blot with CLASP-2 DNA markers to yield unique bands for identifying the individual.

As described above, upon sequencing of numerous independent cDNA products, single nucleotide polymorphisms (SNPs) have been discovered within CLASP-2. These alterations and differences are presented in FIG 11B. They represent mis-sense alterations.

5 If it is determined that certain SNPs are deleterious or advantageous, SNPs can be used as a diagnostic tool through SNP mapping or direct sequencing of the SNP region to determine which isoform is expressed. Additionally, the SNPs can be used as a general SNP marker for chromosomal defects such as rearrangement and translocations.

10 CLASP-2 polynucleotides can be also be used as polymorphic markers for forensic analysis. See generally National Research Council, The Evaluation of Forensic DNA Evidence (Eds. 1996, Pollard *et al.*, National Academy Press, Washington D.C.). The capacity to identify a distinguishing or unique set of forensic markers in an individual is useful for forensic analysis. For example, one can determine whether a blood sample from a suspect matches a blood or other tissue sample from a crime scene by determining whether the set of polymorphic forms occupying selected polymorphic sites is the same in the suspect and the sample. If the set of polymorphic markers does not match between a suspect and a sample, it can be concluded (barring experimental error) that the suspect was not the source of the sample. If the set of markers does match, one can conclude that the DNA from the suspect is consistent with that found at the crime scene. If frequencies of the polymorphic forms at the loci tested have been determined (*e.g.*, by analysis of a suitable population of individuals), one can perform a statistical analysis to determine the probability that a match of suspect and crime scene sample would occur by chance.

25 To make such an identification, PCR technology can be used to amplify DNA sequences taken from very small biological samples such as tissues, *e.g.*, hair or skin, or body fluids, *e.g.*, blood, saliva, or semen found at a crime scene. The amplified sequence can then be compared to a standard, thereby allowing identification of the origin of the biological sample. The CLASP-2 polynucleotide sequences of the present invention can be used to provide polynucleotide reagents, *e.g.*, PCR primers, targeted to specific loci in the human genome, which can enhance the reliability of DNA-based forensic identifications by, for example, providing another "identification marker" (*i.e.* another DNA sequence that is unique to a particular individual). As mentioned above, actual base sequence information can be used for identification as an accurate alternative to patterns formed by restriction enzyme generated fragments. Sequences targeted to noncoding regions of SEQ ID NO: 1, 3, 5 or 9 are particularly appropriate for this use as greater numbers of polymorphisms occur in the

noncoding regions, making it easier to differentiate individuals using this technique.

Examples of polynucleotide reagents include the CLASP-2 nucleotide sequences or portions thereof, *e.g.*, fragments derived from the noncoding regions of SEQ ID NO: 1, 3, 5, or 9 having a length of at least 20 bases, preferably at least 25 bases, and more preferably at least 30 bases.

CLASP-2 polynucleotides can also be used as reagents for paternity testing. The object of paternity testing is usually to determine whether a male is the father of a child. In most cases, the mother of the child is known and thus, the mother's contribution to the child's genotype can be traced. Paternity testing investigates whether the part of the child's genotype not attributable to the mother is consistent with that of the putative father. Paternity testing can be performed by analyzing sets of polymorphisms in the putative father and the child. Of course, the present invention can be expanded to the use of this procedure to determine if one individual is related to another. Even more broadly, the present invention can be employed to determine how related one individual is to another, for example, between races or species.

Bacterial infections are a major cause of health-related problems. However, the emergence of drug resistant bacteria is compromising the therapeutic value of the present spectrum of antibiotics. All the currently used antibiotics are small organic molecules, with certain level of structural similarity. This provides an advantage for bacteria to develop drug resistance, since they need to modify a limited number of genes in order to become resistant to a wide variety of antibiotics. The development of antibiotics with different chemical structure and targets can overcome antibiotic resistance, and provide therapeutic superiority in preventing infection by bacterial pathogens. Additionally, most antibiotics are not naturally occurring compounds and cause minor or sometimes serious side effects. For example, antibiotics used to treat TB can cause hearing loss.

The present invention provides new antibacterial agents. Certain CLASP-2 DNA sequences were difficult to clone and subclone (*see* Example 1). Bacteria harboring certain pieces of CLASP cDNA products were unable to be isolated, indicating that introduction of CLASP sequences compromised bacterial viability. There can be at least two possible reasons why the CLASP cDNA were unable to be cloned, which can reflect a variation of the well-established Modification and Restriction systems found in bacteria (reviewed in Wilson and Murray. (1991) *Annu. Rev. Genet.* 25:585-627; Bickle and Kruger (1993) *Microbiol. Rev.* 57:29-67). This well-described system is used by bacteria to prevent deleterious effects caused by the introduction of foreign DNA. Bacteria can recognize

foreign DNA since it does not have the same modifications (e.g. methylation) as the native DNA. After recognition, the bacteria then digest and eliminate the foreign DNA (restriction). In the first scenario, the CLASP cDNA can be recognized as foreign DNA, and digested and eliminated as in the Modification and Restriction system. However, this would be unique for CLASP cDNA since the bacteria used for cloning cDNA are compromised in the Modification and Restriction system, which makes cloning of cDNA into bacteria a practice common in the art. If this is the case, the bacterial apparatus that specifically recognizes or eliminates CLASP cDNA can provide a novel target to develop antimicrobial agents. The CLASP DNA sequence would be useful in targeting the apparatus as well as an entry point for designing screens to identify potential targets. The second possibility is that CLASP cDNA behaves as an antimicrobial agent (i.e., antibiotic), and prevents bacterial growth. This, in effect, would create a new type of antibiotic mediated by the presence of foreign DNA (i.e. CLASP cDNA). In the case for the CLASP cDNA, the bacteria can recognize the DNA but instead of digesting and eliminating the DNA, the CLASP cDNA can cause a variation of the restriction and prevent the bacteria from growing, imposing a bacteriacidal effect upon the bacteria.

DNA as an antimicrobial agent has significant advantages over currently available agents. First, it is structurally unrelated to any existing antibiotics, and can overcome the present growing drug-resistance problem to structurally common agents. Second, since DNA antimicrobials composed of naturally-occurring human DNA, are expected to have minimal side effects and immune rejection. Third, DNA sequences can be tailored with sequence variation and numerous chemical modifications to circumvent the problem of resistance. Fourth, the antimicrobial DNA can be delivered specifically to bacterial cells through the use of bacteriophages (i.e., bacterial virus) which specifically infect bacteria and do not infect human cells. Further specificity can be generated to infect certain bacteria and bacterial subpopulations. Finally, this system can be economically robust since the generation of DNA and delivery vehicles are inexpensive.

#### **5.5. Polypeptides Encoded by the CLASP-2 Gene Coding Sequence**

In accordance with the invention, a CLASP-2 polynucleotide which encodes the CLASP-2 polypeptides, mutant polypeptides, peptide fragments, CLASP-2 fusion proteins or functional equivalents thereof, can be used to express CLASP-2 proteins in appropriate host cells. In various embodiments, the CLASP-2 polypeptides expressed will be identical or substantially similar to SEQ ID NOs: 2, 4, 6 or 10 or a fragment thereof.

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In some embodiments, altered DNA sequences which can be used in accordance with the invention include deletions, additions or substitutions of different nucleotide residues resulting in a sequence that encodes the same or a functionally equivalent gene product. For example, due to the inherent degeneracy of the genetic code, other DNA sequences which encode substantially the same or a functionally equivalent amino acid sequence, can be used in the practice of the invention for the expression of the CLASP-2 protein. Because of the degeneracy of the genetic code, a large number of functionally identical nucleic acids encode any given protein. For instance, the codons GCA, GCC, GCG and GCU all encode the amino acid alanine. Thus, at every position where an alanine is specified by a codon, the codon can be altered to any of the corresponding codons described without altering the encoded polypeptide. Such nucleic acid variations are "silent variations," which are one species of conservatively modified variations. One of skill will recognize that each codon in a nucleic acid sequence such as SEQ ID NO: 1 (except AUG, which is ordinarily the only codon for methionine, and TGG, which is ordinarily the only codon for tryptophan) can be modified to yield a functionally identical molecule. Accordingly, each silent variation of a nucleic acid which encodes a polypeptide is implicit in each described sequence. Thus, for example, due to the degeneracy of the genetic code, a polypeptide having the sequence of SEQ ID NO: 2 or a fragment thereof, can be encoded by numerous polynucleotides other than SEQ ID NO: 1. Typically, the degenerate sequence will hybridize with SEQ ID NO: 1 under high or moderate stringency conditions, but this is not strictly required (e.g., when a copy of a nucleic acid is created using the maximum codon degeneracy permitted by the genetic code. In such cases, the nucleic acids typically hybridize under moderately stringent hybridization conditions.)

The gene product itself can contain deletions, additions or substitutions of amino acid residues within a CLASP-2 sequence, which result in a silent change thus producing a functionally equivalent CLASP-2 protein. Such conservative amino acid substitutions can be made on the basis of similarity in polarity, charge, solubility, hydrophobicity, hydrophilicity, and/or the amphipathic nature of the residues involved. For example, negatively charged amino acids include aspartic acid and glutamic acid; positively charged amino acids include lysine, histidine and arginine; amino acids with uncharged polar head groups having similar hydrophilicity values include the following: glycine, asparagine, glutamine, serine, threonine, tyrosine; and amino acids with nonpolar head groups include alanine, valine, isoleucine, leucine, phenylalanine, proline, methionine, tryptophan. Creighton, 1984, PROTEINS, has grouped amino acids that are conservative substitutions for



one another as follows: (1) Alanine (A), Glycine (G); (2) Aspartic acid (D), Glutamic acid (E); (3) Asparagine (N), Glutamine (Q); (4) Arginine (R), Lysine (K); (5) Isoleucine (I), Leucine (L), Methionine (M), Valine (V); (6) Phenylalanine (F), Tyrosine (Y), Tryptophan (W); (7) Serine (S), Threonine (T); and (8) Cysteine (C), Methionine (M).

5           The DNA sequences of the invention can be engineered in order to alter a CLASP-2 coding sequence for a variety of ends, including but not limited to, alterations which modify processing and expression of the gene product. For example, mutations can be introduced using techniques which are well known in the art, *e.g.*, site-directed mutagenesis, to insert new restriction sites, to alter glycosylation patterns, phosphorylation, and the like.

10       Based on the domain organization of the CLASP-2 proteins, a large number of CLASP-2 mutant polypeptides can be constructed by modifying or rearranging the nucleotide sequences that encode the CLASP-2 extracellular, transmembrane and cytoplasmic domains.

15           In various embodiments, the present invention provides homologues of the CLASP-2 polypeptides which function as either an CLASP-2 agonists or an CLASP-2 antagonist. In a preferred embodiment, the CLASP-2 agonists and antagonists stimulate or inhibit, respectively, a subset of the biological activities of the naturally occurring form of the CLASP-2 polypeptide. Thus, specific biological effects can be elicited by treatment with a homologue of limited function. In one embodiment, treatment of a subject with a homologue having a subset of the biological activities of the naturally occurring form of the polypeptide  
20       has fewer side effects in a subject relative to treatment with the naturally occurring form of the CLASP-2 polypeptide.

25           The invention contemplates both full-length CLASP-2 polypeptides and fragments, *e.g.*, fragments having a length of at least about 10, often 20, frequently 50 or 100 residues substantially identical to the exemplified CLASP-2 polypeptide sequences of the invention. Protein fragments can be “free-standing,” or comprised within a larger polypeptide of which the fragment forms a part or region, most preferably as a single continuous region. Representative examples of polypeptide fragments of the invention, include, for example, fragments from about amino acid number 1-20, 2 1-40, 4 1-60, 61-80, 81-100, 102-120, 121-140, 141-160, 161-180, 181-200, or 201 to the end of the coding  
30       region. Moreover, polypeptide fragments can be about 20, 30, 40, 50, 60, 70, 80, 90, 100, 110, 120, 130, 140, 150, 200 amino acids in length. In this context “about” includes the particularly recited ranges, larger or smaller by several (5, 4, 3, 2, or 1) amino acids, at either extreme or at both extremes.

Preferred polypeptide fragments include the CLASP-2 protein. Further preferred polypeptide fragments include the CLASP-2 protein having a continuous series of deleted residues from the amino or the carboxy terminus, or both. For example, any number of amino acids, ranging from 1-X, can be deleted from the amino terminus of either the CLASP-2 polypeptide. Furthermore, any combination of the above amino and carboxy terminus deletions are preferred. Similarly, polynucleotide fragments encoding these CLASP-2 polypeptide fragments are also preferred.

Even if deletion of one or more amino acids from the N-terminus of a protein results in modification or loss of one or more biological functions of the protein, other biological activities can still be retained. Thus, the ability of shortened CLASP-2 muteins to induce and/or bind to antibodies which recognize the complete or mature forms of the polypeptides generally will be retained when less than the majority of the residues of the complete or mature polypeptide are removed from the N-terminus. Whether a particular polypeptide lacking N-terminal residues of a complete polypeptide retains such immunologic activities can readily be determined by routine methods described herein and otherwise known in the art. It is not unlikely that a CLASP-2 mutein with a large number of deleted N-terminal amino acid residues can retain some biological or immunogenic activities. In fact, peptides composed of as few as four CLASP-2 amino acid residues can often evoke an immune response.

Homologues of the CLASP-2 polypeptide can be generated by mutagenesis, *e.g.*, discrete point mutation or truncation of the CLASP-2 polypeptide. As used herein, the term "homologue" refers to a variant form of the CLASP-2 polypeptide which acts as an agonist or antagonist of the activity of the CLASP-2 polypeptide. An agonist of the CLASP-2 polypeptide can retain substantially the same, or a subset, of the biological activities of the CLASP-2 polypeptide. An antagonist of the CLASP-2 polypeptide can inhibit one or more of the activities of the naturally occurring form of the CLASP-2 polypeptide, by, for example, competitively binding to a downstream or upstream member of the CLASP-2 molecular pathway which includes the CLASP-2 polypeptide.

Modulation can be assayed by determining any parameter that is indirectly or directly affected by the expression of the target gene. Such parameters include, *e.g.*, changes in RNA or protein levels, changes in protein activity, changes in product levels, changes in downstream gene expression, changes in reporter gene transcription (luciferase, CAT,  $\beta$ -galactosidase,  $\beta$ -glucuronidase, GFP (*see, e.g.*, Mistili & Spector, 1997, *Nature Biotechnology* 15: 961-964); changes in signal transduction, phosphorylation and

dephosphorylation, receptor-ligand interactions, second messenger concentrations (e.g., cGMP, cAMP, IP<sub>3</sub>, and Ca<sup>2+</sup>), and cell growth. These assays can be *in vitro*, *in vivo*, and *ex vivo*. Such functional effects can be measured by any means known to those skilled in the art, e.g., measurement of RNA or protein levels, measurement of RNA stability, identification of downstream or reporter gene expression, e.g., via chemiluminescence, fluorescence, colorimetric reactions, antibody binding, inducible markers, ligand binding assays; changes in intracellular second messengers such as cGMP and inositol triphosphate (IP<sub>3</sub>); changes in intracellular calcium levels; cytokine release, and the like.

#### **5.5.1. Synthesis or Expression of CLASP-2 Polypeptide Expression Systems**

In order to express a biologically active CLASP-2, the nucleotide sequence coding for CLASP-2, or a functional equivalent, is inserted into an appropriate expression vector. The CLASP-2 gene product as well as host cells or cell lines transfected or transformed with recombinant CLASP-2 expression vectors can be used for a variety of purposes. These include, but are not limited to, generating antibodies (*i.e.*, monoclonal or polyclonal) that competitively inhibit activity of CLASP-2 protein and neutralize its activity; antibodies that activate CLASP-2 function and antibodies that detect its presence on the cell surface or in solution. Anti-CLASP-2 antibodies can be used in detecting and quantifying expression of CLASP-2 levels in cells and tissues such as lymphocytes and macrophages, as well as isolating CLASP-2-positive cells from a cell mixture.

Methods which are well known to those skilled in the art can be used to construct recombinant expression vectors containing the CLASP-2 coding sequence and appropriate transcriptional/translational control signals. These methods include *in vitro* recombinant DNA techniques, synthetic techniques and *in vivo* recombination/genetic recombination. (See, e.g., the techniques described in Sambrook *et al.*, 1989, Molecular Cloning A Laboratory Manual, Cold Spring Harbor Laboratory, N.Y. and Ausubel *et al.*, *supra*). The recombinant expression vectors of the invention comprise a nucleic acid of the invention in a form suitable for expression of the nucleic acid in a host cell, which means that the recombinant expression vectors include one or more regulatory sequences, selected on the basis of the host cells to be used for expression, which is operatively linked to the nucleic acid sequence to be expressed. It will be appreciated by those skilled in the art that the design of the expression vector can depend on such factors as the choice of the host cell to be transformed, the level of expression of polypeptide desired, and the like. The expression vectors of the invention can be introduced into host cells to thereby produce polypeptides or

peptides, including fusion polypeptides or peptides, encoded by nucleic acids as described herein (*e.g.*, CLASP-2 polypeptides, mutant forms of CLASP-2, fusion polypeptides, and the like).

A variety of host-expression vector systems can be utilized to express a CLASP-2 coding sequence. These include, but are not limited to, microorganisms such as bacteria transformed with recombinant bacteriophage DNA, plasmid DNA, or cosmid DNA expression vectors containing the CLASP-2 coding sequence; yeast transformed with recombinant yeast expression vectors containing the CLASP-2 coding sequence; insect cell systems infected with recombinant virus expression vectors (*e.g.*, baculovirus) containing the CLASP-2 coding sequence; plant cell systems infected with recombinant virus expression vectors (*e.g.*, cauliflower mosaic virus, CaMV; tobacco mosaic virus, TMV) or transformed with recombinant plasmid expression vectors (*e.g.*, Ti plasmid) containing the CLASP-2 coding sequence; or animal cell systems. The expression elements of these systems vary in their strength and specificities. Depending on the host/vector system utilized, any of a number of suitable transcription and translation elements, including constitutive and inducible promoters, can be used in the expression vector. For example, when cloning in bacterial systems, inducible promoters such as pL of bacteriophage  $\lambda$ , plac, ptrp, ptac (ptrp-lac hybrid promoter; cytomegalovirus promoter) and the like can be used; when cloning in insect cell systems, promoters such as the baculovirus polyhedron promoter can be used; when cloning in plant cell systems, promoters derived from the genome of plant cells (*e.g.*, heat shock promoters; the promoter for the small subunit of RUBISCO; the promoter for the chlorophyll  $\alpha/\beta$  binding protein) or from plant viruses (*e.g.*, the <sup>35</sup>S RNA promoter of CaMV; the coat protein promoter of TMV) can be used; when cloning in mammalian cell systems, promoters derived from the genome of mammalian cells (*e.g.*, metallothionein promoter) or from mammalian viruses (*e.g.*, the adenovirus late promoter; the vaccinia virus 7.5K promoter) can be used; when generating cell lines that contain multiple copies of the CLASP-2 DNA, SV40-, BPV- and EBV-based vectors can be used with an appropriate selectable marker.

In bacterial systems a number of expression vectors can be advantageously selected depending upon the use intended for the expressed CLASP-2 product. For example, when large quantities of CLASP-2 protein are to be produced for the generation of antibodies or to screen peptide libraries, vectors which direct the expression of high levels of fusion protein products that are readily purified can be desirable. Such vectors include, but are not

limited to, the *E. coli* expression vector pUR278 (Ruther *et al.*, 1983, EMBO J. 2: 1791), in which the CLASP-2 coding sequence can be ligated into the vector in frame with the lacZ coding region so that a hybrid protein is produced; pIN vectors (Inouye & Inouye, 1985, Nucleic acids Res. 13: 3101-3109; Van Heeke & Schuster, 1989, J. Biol. Chem. 264: 5503-5509); and the like. pGEX vectors may also be used to express foreign polypeptides as fusion proteins with glutathione S-transferase (GST). In general, such fusion proteins are soluble and can easily be purified from lysed cells by adsorption to glutathione-agarose beads followed by elution in the presence of free glutathione. Proteins made in such systems may be designed to include heparin, thrombin, or factor XA protease cleavage sites so that the cloned polypeptide of interest can be released from the GST moiety at will.

In yeast, a number of vectors containing constitutive or inducible promoters can be used. (Current Protocols in Molecular Biology, Vol. 2, 1988 (Suppl. 1999), Ed. Ausubel *et al.*, Greene Publish. Assoc. & Wiley Interscience, Ch. 13; Grant *et al.*, 1987, Expression and Secretion Vectors for Yeast, in Methods in Enzymology, Eds. Wu & Grossman, 1987, Acad. Press, N.Y., Vol. 153, pp. 516-544; Glover, 1986, DNA Cloning, Vol. II, IRL Press, Wash., D.C., Ch. 3; and Bitter, 1987, Heterologous Gene Expression in Yeast, Methods in Enzymology, Eds. Berger & Kimmel, Acad. Press, N.Y., Vol. 152, pp. 673-684; and The Molecular Biology of the Yeast *Saccharomyces*, 1982, Eds. Strathern *et al.*, Cold Spring Harbor Press, Vols. I and II.)

In cases where plant expression vectors are used, the expression of the CLASP-2 coding sequence can be driven by any of a number of promoters. For example, viral promoters such as the 35S RNA and 19S RNA promoters of CaMV (Brisson *et al.*, 1984, Nature 310: 511-514), or the coat protein promoter of TMV (Takamatsu *et al.*, 1987, EMBO J. 6: 307-311) can be used; alternatively, plant promoters such as the small subunit of RUBISCO (Coruzzi *et al.*, 1984, EMBO J. 3: 1671-1680; Broglie *et al.*, 1984, Science 224: 838-843); or heat shock promoters, *e.g.*, soybean hsp17.5-E or hsp17.3-B (Gurley *et al.*, 1986, Mol. Cell. Biol. 6: 559-565) can be used. These constructs can be introduced into plant cells using Ti plasmids, Ri plasmids, plant virus vectors, direct DNA transformation, microinjection, electroporation, and the like. (Weissbach & Weissbach, 1988, Methods for Plant Molecular Biology, Academic Press, NY, Section VIII, pp. 421-463; and Grierson & Corey, 1988, Plant Molecular Biology, 2d Ed., Blackie, London, Ch. 7-9.)

An alternative expression system which could be used to express CLASP-2 is an insect system. In one such system, *Autographa californica* nuclear polyhedrosis virus (AcNPV) is used as a vector to express foreign genes. The virus grows in *Spodoptera*



*frugiperda* cells. The CLASP-2 coding sequence can be cloned into non-essential regions (e.g., the polyhedron gene) of the virus and placed under control of an AcNPV promoter (e.g., the polyhedron promoter). Successful insertion of the CLASP-2 coding sequence will result in inactivation of the polyhedron gene and production of non-occluded recombinant virus (i.e., virus lacking the proteinaceous coat coded for by the polyhedron gene). These recombinant viruses are then used to infect *Spodoptera frugiperda* cells in which the inserted gene is expressed. (see, e.g., Smith *et al.*, 1983, J. Virol. 46: 584; Smith, U.S. Patent No. 4,215,051).

In mammalian host cells, a number of viral based expression systems can be utilized. In cases where an adenovirus is used as an expression vector, the CLASP-2 coding sequence can be ligated to an adenovirus transcription/translation control complex, e.g., the late promoter and tripartite leader sequence. This chimeric gene can then be inserted in the adenovirus genome by *in vitro* or *in vivo* recombination. Insertion in a non-essential region of the viral genome (e.g., region E1 or E3) will result in a recombinant virus that is viable and capable of expressing CLASP-2 in infected hosts. (See, e.g., Logan & Shenk, 1984, Proc. Natl. Acad. Sci. U.S.A. 81: 3655-3659). Alternatively, the vaccinia 7.5K promoter can be used. (See, e.g., Mackett *et al.*, 1982, Proc. Natl. Acad. Sci. U.S.A. 79: 7415-7419; Mackett *et al.*, 1984, J. Virol. 49: 857-864; Panicali *et al.*, 1982, Proc. Natl. Acad. Sci. U.S.A. 79: 4927-4931). Regulatable expression vectors such as the tetracycline repressible vectors can also be used to express a coding sequence in a controlled fashion.

Specific initiation signals can also be required for efficient translation of inserted CLASP-2 coding sequences. These signals include the ATG initiation codon and adjacent sequences. In cases where the entire CLASP-2 gene, including its own initiation codon and adjacent sequences, is inserted into the appropriate expression vector, no additional translational control signals can be needed. However, in cases where only a portion of the CLASP-2 coding sequence is inserted, exogenous translational control signals, including the ATG initiation codon, must be provided. Furthermore, the initiation codon must be in phase with the reading frame of the CLASP-2 coding sequence to ensure translation of the entire insert. These exogenous translational control signals and initiation codons can be of a variety of origins, both natural and synthetic. The efficiency of expression can be enhanced by the inclusion of appropriate transcription enhancer elements, transcription terminators, and the like. (see Bittner *et al.*, 1987, Methods in Enzymol. 153: 516-544).

5 In addition, a host cell strain can be chosen which modulates the expression of  
the inserted sequences, or modifies and processes the gene product in a specific fashion  
desired. Such modifications (*e.g.*, glycosylation) and processing (*e.g.*, cleavage) of protein  
products can be important for the function of the protein. The presence of several consensus  
10 N-glycosylation sites in CLASP-2 extracellular domains support the possibility that proper  
modification can play a role in CLASP-2 function. Different host cells have characteristic  
and specific mechanisms for the post-translational processing and modification of proteins.  
Appropriate cell lines or host systems can be chosen to ensure the correct modification and  
processing of the foreign protein expressed. To this end, eukaryotic host cells which possess  
15 the cellular machinery for proper processing of the primary transcript, glycosylation, and  
phosphorylation of the gene product can be used. Such mammalian host cells include, but are  
not limited to, CHO, VERO, BHK, HeLa, COS, MDCK, 293, WI38, and the like.

Host cells transformed with nucleotide sequences encoding CLASP-2 may be  
cultured under conditions suitable for the expression and recovery of the soluble protein from  
cell culture. The protein produced by a transformed cell may be secreted or contained  
15 intracellularly depending on the sequence and/or the vector used. As will be understood by  
those of skill in the art, expression vectors containing polynucleotides which encode CLASP-  
2 may be designed to contain signal sequences which direct secretion of CLASP-2 through a  
prokaryotic or eukaryotic cell membrane. Other constructions may be used to join sequences  
20 encoding CLASP-2 to nucleotide sequence encoding a polypeptide domain which will  
facilitate purification of soluble proteins. Such purification facilitating domains include, but  
are not limited to, metal chelating peptides such as histidine-tryptophan modules that allow  
purification on immobilized metals, protein A domains that allow purification on  
immobilized immunoglobulin,

25 For long-term, high-yield production of recombinant proteins, stable  
expression is preferred. For example, cell lines which stably express CLASP-2 proteins can  
be engineered. Rather than using expression vectors which contain viral origins of  
replication, host cells can be transformed with the CLASP-2 DNA controlled by appropriate  
expression control elements (*e.g.*, promoter, enhancer, sequences, transcription terminators,  
30 polyadenylation sites, and the like.), and a selectable marker. Following the introduction of  
foreign DNA, engineered cells can be allowed to grow for 1-2 days in an enriched medium,  
and then switched to a selective medium. The selectable marker in the recombinant plasmid  
confers resistance to the selection and allows cells to stably integrate the plasmid into their  
chromosomes and grow to form foci which in turn can be cloned and expanded into cell lines.

This method can advantageously be used to engineer cell lines which express the CLASP-2 protein(s) on the cell surface. Such engineered cell lines are particularly useful in screening for molecules or drugs that affect CLASP-2 function.

A number of selection systems can be used, including but not limited to, the herpes simplex virus thymidine kinase (Wigler *et al.*, 1977, Cell 11: 223), hypoxanthine-guanine phosphoribosyltransferase (Szybalska & Szybalski, 1962, Proc. Natl. Acad. Sci. U.S.A. 48: 2026), and adenine phosphoribosyltransferase (Lowy *et al.*, 1980, Cell 22: 817) genes which can be employed in tk<sup>-</sup>, hgprt<sup>-</sup> or aprt<sup>-</sup> cells, respectively. Also, antimetabolite resistance can be used as the basis of selection for *dhfr*, which confers resistance to methotrexate (Wigler *et al.*, 1980, Natl. Acad. Sci. U.S.A. 77: 3567; O'Hare *et al.*, 1981, Proc. Natl. Acad. Sci. U.S.A. 78: 1527); *gpt*, which confers resistance to mycophenolic acid (Mulligan & Berg, 1981, Proc. Natl. Acad. Sci. U.S.A. 78: 2072); *neo*, which confers resistance to the aminoglycoside G-418 (Colberre-Garapin *et al.*, 1981, J. Mol. Biol. 150: 1); and *hygro*, which confers resistance to hygromycin (Santerre *et al.*, 1984, Gene 30: 147). Additional selectable genes have been described, namely *trpB*, which allows cells to utilize indole in place of tryptophan; *hisD*, which allows cells to utilize histinol in place of histidine (Hartman & Mulligan, 1988, Proc. Natl. Acad. Sci. U.S.A. 85: 8047); *ODC* (ornithine decarboxylase) which confers resistance to the ornithine decarboxylase inhibitor, 2-(difluoromethyl)-DL-ornithine, DFMO (McConlogue L., 1987, In: Current Communications in Molecular Biology, Cold Spring Harbor Laboratory ed.) and glutamine synthetase (Bebbington *et al.*, 1992, Biotech 10: 169).

In an alternate embodiment of the invention, the coding sequence of CLASP-2 could be synthesized in whole or in part, using chemical methods well known in the art. (See, *e.g.*, Caruthers *et al.*, 1980, Nuc. Acids Res. Symp. Ser. 7: 215-233; Crea and Horn, 180, Nuc. Acids Res. 9(10): 2331; Matteucci and Caruthers, 1980, Tetrahedron Letter 21: 719; and Chow and Kempe, 1981, Nuc. Acids Res. 9(12): 2807-2817.) Alternatively, the protein itself could be produced using chemical methods to synthesize a CLASP-2 amino acid sequence in whole or in part. For example, peptides can be synthesized by solid phase techniques, cleaved from the resin, and purified by preparative high performance liquid chromatography. (See Creighton, 1983, Proteins Structures And Molecular Principles, W.H. Freeman and Co., N.Y. pp. 50-60). The composition of the synthetic polypeptides can be confirmed by amino acid analysis or sequencing (*e.g.*, the Edman degradation procedure; see Creighton, 1983, Proteins, Structures and Molecular Principles, W.H. Freeman and Co., N.Y., pp. 34-49).

In some embodiments, the CLASP-2 polypeptide contains non-naturally occurring amino acids or amino acid analogs (*i.e.*, compounds that have the same basic chemical structure as a naturally occurring amino acid, *i.e.*, an  $\alpha$  carbon that is bound to a hydrogen, a carboxyl group, an amino group, and an R group, *e.g.*, homoserine, norleucine, methionine sulfoxide, methionine methyl sulfonium).

### **5.5.2. Identification of Cells That Express CLASP-2**

The recombinant host cells which contain the coding sequence and which express a CLASP-2 gene product or fragments thereof can be identified by at least four general approaches; (a) DNA-DNA or DNA-RNA hybridization; (b) the presence or absence of “marker” gene functions; (c) assessing the level of transcription as measured by the expression of CLASP-2 mRNA transcripts in the host cell; and (d) detection of the gene product as measured by immunoassay or by its biological activity. Prior to the identification of gene expression, the host cells can be first mutagenized in an effort to increase the level of expression of CLASP-2, especially in cell lines that produce low amounts of CLASP-2.

In the first approach, the presence of the CLASP-2 coding sequence inserted in the expression vector can be detected by DNA-DNA or DNA-RNA hybridization using probes comprising nucleotide sequences that are homologous to the CLASP-2 coding sequence, respectively, or portions or derivatives thereof.

In the second approach, the recombinant expression vector/host system can be identified and selected based upon the presence or absence of certain “marker” gene functions (*e.g.*, thymidine kinase activity, resistance to antibiotics, resistance to methotrexate, transformation phenotype, occlusion body formation in baculovirus, and the like). For example, if the CLASP-2 coding sequence is inserted within a marker gene sequence of the vector, recombinants containing the CLASP-2 coding sequence can be identified by the absence of the marker gene function. Alternatively, a marker gene can be placed in tandem with the CLASP-2 sequence under the control of the same or different promoter used to control the expression of the CLASP-2 coding sequence. Expression of the marker in response to induction or selection indicates expression of the CLASP-2 coding sequence.

In the third approach, transcriptional activity for the CLASP-2 coding region can be assessed by hybridization assays. For example, RNA can be isolated and analyzed by Northern blot using a probe homologous to the CLASP-2 coding sequence or particular portions thereof. Alternatively, total nucleic acids of the host cell can be extracted and

assayed for hybridization to such probes. Additionally, reverse transcription-polymerase chain reactions can be used to detect low levels of gene expression.

In the fourth approach, the expression of the CLASP-2 protein product can be assessed immunologically, for example by Western blots, immunoassays such as radioimmuno-precipitation, enzyme-linked immunoassays, fluorescent activated cell sorting (“FACS”), and the like. This can be achieved by using an anti-CLASP-2 antibody. Alternatively, CLASP-2 protein can be expressed as a fusion protein with green-fluorescent protein to facilitate its detection in cells (United States Patent Nos. 5,491,084; 5,804,387; 5,777,079).

Identification of cells or tissues expressing CLASP protein or mRNA, especially CLASP-2 isoforms, can be useful for determining normal and abnormal CLASP expression in a given cell or tissue. As discussed above, a number of CLASP-2 isoforms have been identified, *e.g.*, in Jurkat cells, peripheral blood, and brain. The identification of mRNA or protein expression in various cell types and tissues can allow for identification of isoforms improperly expressed in either a spatial or temporal manner. Expression of hCLASP-2D isoform in hematopoietic cells may cause problems due to the presence of the SH3 domain not seen in the Jurkat and peripheral blood isoforms.

Other molecules in the immune system may also interact with portions of hCLASP2D. However, the absence of the PBM domain in the hCLASP-2D isoform may be necessary for function in certain cell types or tissues. Similarly, expression of CLASP isoforms 2A, 2B, and 2C in brain may cause problems for different reasons: the PBM present in these isoforms may interfere with a particular function by binding any of the known PDZ domain protein involved in formation of the neurological synapse. Similarly, the lack of an SH3 domain may cause an inappropriate response due to interactions with only a subset of molecules required for CLASP-2 function in the brain.

### **5.5.3. Uses of CLASP-2 Engineered Host Cells**

In one embodiment of the invention, the CLASP-2 protein and/or cell lines that express CLASP-2 can be used to screen for antibodies, peptides, small molecules, natural and synthetic compounds or other cell bound or soluble molecules that bind to the CLASP-2 protein resulting in stimulation or inhibition of CLASP-2 function. For example, anti-CLASP-2 antibodies can be used to inhibit or stimulate CLASP-2 function and to detect its presence. Alternatively, screening of peptide libraries with recombinantly expressed soluble CLASP-2 protein or cell lines expressing CLASP-2 protein can be useful for identification of



therapeutic molecules that function by inhibiting or stimulating the biological activity of CLASP-2. The uses of the CLASP-2 protein and engineered cell lines, described in the subsections below, can be employed equally well for homologous CLASP-2 genes in various species.

5 In a specific embodiment of the invention, cell lines may be engineered to express the extracellular or intracellular domain of CLASP fused to another molecule such as GST. In addition, CLASP, its extracellular domain or its intracellular domain may be fused to an immunoglobulin constant region (Hollenbaugh and Aruffo, 1992, Current Protocols in Immunology, Unit 10.19; Aruffo *et al.*, 1990, Cell 61: 1303) to produce a soluble molecule  
10 with increased half life. The soluble protein or fusion protein can be used in binding assays, affinity chromatography, immunoprecipitation, Western blot, and the like. Synthetic compounds, natural products, and other sources of potentially biologically active materials can be screened in assays that are well known in the art.

15 Random peptide libraries consisting of all possible combinations of amino acids attached to a solid phase support can be used to identify peptides that are able to bind to a specific domain of CLASP-2 (Lam, K.S. *et al.*, 1991, Nature 354: 82-84). The screening of peptide libraries can have therapeutic value in the discovery of pharmaceutical agents that stimulate or inhibit the biological activity of CLASP-2.

20 Identification of molecules that are able to bind to the CLASP-2 protein can be accomplished by screening a peptide library with recombinant soluble CLASP-2 protein. Methods for expression and purification of CLASP-2 are described in Section 5.7, *supra*, and can be used to express recombinant full length CLASP-2 or fragments of CLASP-2 depending on the functional domains of interest. Such domains include CLASP-2 extracellular domain, transmembrane domain, CLASP-2 intracellular domain, ITAM  
25 containing domain, tyrosine phosphorylation site containing domain, cysteine cluster containing domain, cadherin motif containing domain, and coil/coil domain.

To identify and isolate the peptide/solid phase support that interacts and forms a complex with CLASP-2, it is necessary to label or "tag" the CLASP-2 molecule. The CLASP-2 protein can be conjugated to enzymes such as alkaline phosphatase or horseradish  
30 peroxidase or to other reagents such as fluorescent labels which can include fluorescein isothiocyanate (FITC), phycoerythrin (PE) or rhodamine. Conjugation of any given label to CLASP-2 can be performed using techniques that are well known in the art. Alternatively, CLASP-2 expression vectors can be engineered to express a chimeric CLASP-2 protein containing an epitope for which a commercially available antibody exist. The epitope-

specific antibody can be tagged with a detectable label using methods well known in the art including an enzyme, a fluorescent dye or colored or magnetic beads.

The “tagged” CLASP-2 conjugate is incubated with the random peptide library for 30 minutes to one hour at 22°C to allow complex formation between CLASP-2 and peptide species within the library. The library is then washed to remove any unbound protein. If CLASP-2 has been conjugated to alkaline phosphatase or horseradish peroxidase the whole library is poured into a petri dish containing substrates for either alkaline phosphatase or peroxidase, for example, 5-bromo-4-chloro-3-indoyl phosphate (BCIP) or 3,3',4,4"-diaminobenzidine (DAB), respectively. After incubating for several minutes, the peptide/solid phase- CLASP-2 complex changes color, and can be easily identified and isolated physically under a dissecting microscope with a micromanipulator. If a fluorescent tagged CLASP-2 molecule has been used, complexes can be isolated by fluorescence activated sorting. If a chimeric CLASP-2 protein expressing a heterologous epitope has been used, detection of the peptide/CLASP-2 complex can be accomplished by using a labeled epitope-specific antibody. Once isolated, the identity of the peptide attached to the solid phase support can be determined by peptide sequencing.

In addition to using soluble CLASP-2 molecules, in another embodiment, it is possible to detect peptides that bind to cell-associated CLASP-2 using intact cells. The use of intact cells is preferred for use with cell surface molecules. Methods for generating cell lines expressing CLASP-2 are described in Section 5.8. The cells used in this technique can be either live or fixed cells. The cells can be incubated with the random peptide library and bind to certain peptides in the library to form a “rosette” between the target cells and the relevant solid phase support/peptide. The rosette can thereafter be isolated by differential centrifugation or removed physically under a dissecting microscope. Techniques for screening combinatorial libraries are known in the art (Gallop *et al.*, 1994, J. Med. Chem., 37: 1233; Gordon, 1994, J. Med. Chem., 37: 1385).

As an alternative to whole cell assays for membrane bound receptors or receptors that require the lipid domain of the cell membrane to be functional, CLASP-2 molecules can be reconstituted into liposomes where label or “tag” can be attached.

#### **5.5.4. CLASP-2 Fusion Proteins**

In another embodiment of the invention, a CLASP-2 or a modified CLASP-2 sequence can be ligated to a heterologous sequence to encode a fusion protein. For example, for screening of peptide libraries for molecules that bind CLASP-2, it can be useful to

produce a chimeric CLASP-2 protein expressing a heterologous epitope that is recognized by a commercially available antibody. A fusion protein can also be engineered to contain a cleavage site located between a CLASP-2 sequence and the heterologous protein sequence, so that the CLASP-2 can be cleaved away from the heterologous moiety. In one embodiment, fusion proteins of the invention can contain the CLASP-2 extracellular domain comprising at least about residues 1 through 816 or fragment thereof. In another embodiment, fusion proteins can contain the CLASP-2 intracellular domain comprising at least about residue 843 through the end of the CLASP-2 sequence or fragment thereof.

### **5.6. Cloning Alleles, Variants, and Species Homologs of CLASP-2**

In order to clone the full length cDNA sequence from any species encoding a CLASP-2 cDNA, or to clone variant forms of the molecule, labeled DNA probes made from nucleic acid fragments corresponding to any partial cDNA disclosed herein can be used to screen a cDNA library derived from lymphoid cells or brain cells. More specifically, oligonucleotides corresponding to either the 5' or 3' terminus of the cDNA sequence can be used to obtain longer nucleotide sequences. Briefly, the library can be plated out to yield a maximum of 30,000 pfu for each 150 mm plate. Approximately 40 plates can be screened. The plates are incubated at 37°C until the plaques reach a diameter of 0.25 mm or are just beginning to make contact with one another (3-8 hours). Nylon filters are placed onto the soft top agarose and after 60 seconds, the filters are peeled off and floated on a DNA denaturing solution consisting of 0.4N sodium hydroxide. The filters are then immersed in neutralizing solution consisting of 1M Tris-HCl, pH 7.5, before being allowed to air dry. The filters are prehybridized in hybridization buffer such as casein buffer containing 10% dextran sulfate, 0.5M NaCl, 50mM Tris-HCl, pH 7.5, 0.1% sodium pyrophosphate, 1% casein, 1% SDS, and denatured salmon sperm DNA at 0.5 mg/ml for 6 hours at 60°C. The radiolabeled probe is then denatured by heating to 95°C for 2 minutes and then added to the prehybridization solution containing the filters. The filters are hybridized at 60°C for 16 hours. The filters are then washed in 1X wash mix (10X wash mix contains 3M NaCl, 0.6M Tris base, and 0.02M EDTA) twice for 5 minutes each at room temperature, then in 1X wash mix containing 1% SDS at 60°C for 30 minutes, and finally in 0.3X wash mix containing 0.1% SDS at 60°C for 30 minutes. The filters are then air dried and exposed to x-ray film for autoradiography. After developing, the film is aligned with the filters to select a positive plaque. If a single, isolated positive plaque cannot be obtained, the agar plug containing the

plaques will be removed and placed in lambda dilution buffer containing 0.1M NaCl, 0.01M magnesium sulfate, 0.035M Tris HCl, pH 7.5, 0.01% gelatin. The phage can then be replated and rescreened to obtain single, well isolated positive plaques. Positive plaques can be isolated and the cDNA clones sequenced using primers based on the known cDNA sequence.

5 This step can be repeated until a full length cDNA is obtained.

It can be necessary to screen multiple cDNA libraries from different tissues to obtain a full length cDNA. In the event that it is difficult to identify cDNA clones encoding the complete 5' terminal coding region, an often encountered situation in cDNA cloning, the RACE (Rapid Amplification of cDNA Ends) technique can be used. RACE is a proven PCR-  
10 based strategy for amplifying the 5' end of incomplete cDNAs. 5'-RACE-Ready RNA synthesized from human tissues containing a unique anchor sequence is commercially available (Clontech). To obtain the 5' end of the cDNA, PCR is carried out on 5'-RACE-Ready cDNA using the provided anchor primer and the 3' primer. A secondary PCR reaction is then carried out using the anchored primer and a nested 3' primer according to the  
15 manufacturer's instructions. Once obtained, the full length cDNA sequence can be translated into amino acid sequence and examined for certain landmarks such as a continuous open reading frame flanked by translation initiation and termination sites, a cadherin-like domain, an ITAM domain, a tyrosine phosphorylation site, a cysteine cluster, a transmembrane domain, and finally overall structural similarity to the CLASP-2 genes disclosed herein. *See*,  
20 Ponassi *et al.*, 1999, Mech. Dev. 80: 207-212; Isakov, 1998, Receptor Channels 5: 243-253; Borroto *et al.*, 1997, Biopolymers 42: 75-88; Dimitratos *et al.*, 1997, Mech. Dev. 63: 127-130; Apperson *et al.*, 1996, J. Neurosci. 16: 6839-6852; Ozawa *et al.*, 1990, Mech. Dev. 33: 49-56, which discuss protein domains and are incorporated herein by reference.

### **5.7. Modulating Expression of Endogenous CLASP-2 Genes**

25 Alternatively, the expression characteristics of an endogenous CLASP-2 gene within a cell population can be modified by inserting a heterologous DNA regulatory element into the genome of the cell line such that the inserted regulatory element is operatively linked with the endogenous CLASP-2 gene. For example, an endogenous CLASP-2 gene which is normally "transcriptionally silent", *i.e.*, an *CLASP-2* gene which is normally not expressed, or  
30 is expressed only at very low levels in a cell population, can be activated by inserting a regulatory element which is capable of promoting the expression of a normally expressed gene product in the cells. Alternatively, a transcriptionally silent, endogenous CLASP-2 gene

can be activated by insertion of a promiscuous regulatory element that works across cell types.

A heterologous regulatory element can be inserted into a cell line population, such that it is operatively linked with an endogenous CLASP-2 gene, using techniques, such as targeted homologous recombination, which are well known to those of skill in the art, (see e.g., in Chappel, U.S. Patent No. 5,272,071; PCT publication No. WO 91/06667, published Jan 16, 1991).

### **5.8. Anti-CLASP-2 Antibodies**

Various procedures known in the art can be used for the production of antibodies to epitopes of the natural and recombinantly produced CLASP-2 protein. Such antibodies include, but are not limited to, polyclonal, monoclonal, chimeric, single chain, human or humanized, IgG, IgM, IgA, IgD or IgE, a complementarity determining region, Fab fragments, F(ab')<sub>2</sub> and fragments produced by an Fab expression library as well as anti-idiotypic antibodies. Antibodies which compete for CLASP-2 binding are especially preferred for diagnostics and therapeutics.

Monoclonal antibodies that bind CLASP-2 can be radioactively labeled allowing one to follow their location and distribution in the body after injection. Radioisotope tagged antibodies can be used as a non-invasive diagnostic tool for imaging *de novo* lymphoid tumors and metastases that express CLASP-2.

Immunotoxins can also be designed which target cytotoxic agents to specific sites in the body. For example, high affinity CLASP-2 specific monoclonal antibodies can be covalently complexed to bacterial or plant toxins, such as diphtheria toxin or ricin. A general method of preparation of antibody/hybrid molecules can involve use of thiol-crosslinking reagents such as SPDP, which attack the primary amino groups on the antibody and by disulfide exchange, attach the toxin to the antibody. The hybrid antibodies can be used to specifically eliminate CLASP-2 expressing lymphocytes.

For the production of antibodies, various host animals can be immunized by injection with the recombinant or naturally purified CLASP-2 protein, fusion protein or peptides, including but not limited to goats, rabbits, mice, rats, hamsters, and the like

Various adjuvants can be used to increase the immunological response, depending on the host species, including but not limited to Freund's (complete and incomplete), mineral gels such as aluminum hydroxide, surface active substances such as lysolecithin, pluronic polyols, polyanions, peptides, oil emulsions, keyhole limpet hemocyanin, dinitrophenol, and poten-



tially useful human adjuvants such as BCG (bacilli Calmette-Guerin) and *Corynebacterium parvum*.

Monoclonal antibodies to CLASP-2 can be prepared by using any technique which provides for the production of antibody molecules by continuous cell lines in culture. These include, but are not limited to, the hybridoma technique originally described by Kohler and Milstein, (*Nature*, 1975, 256: 495-497), the human B-cell hybridoma technique (Kosbor *et al.*, 1983, *Immunology Today*, 4: 72; Cote *et al.*, 1983, *Proc. Natl. Acad. Sci. U.S.A.*, 80: 2026-2030) and the EBV-hybridoma technique (Cole *et al.*, 1985, *Monoclonal Antibodies and Cancer Therapy*, Alan R. Liss, Inc., pp. 77-96). In addition, techniques developed for the production of "chimeric antibodies" (Morrison *et al.*, 1984, *Proc. Natl. Acad. Sci. U.S.A.*, 81: 6851-6855; Neuberger *et al.*, 1984, *Nature*, 312: 604-608; Takeda *et al.*, 1985, *Nature*, 314: 452-454) by splicing the genes from a mouse antibody molecule of appropriate antigen specificity together with genes from a human antibody molecule of appropriate biological activity can be used. Alternatively, techniques described for the production of single chain antibodies (U.S. Patent 4,946,778) can be adapted to produce CLASP-2 -specific single chain antibodies. In some embodiments, phage display technology is used to identify antibodies and heteromeric Fab fragments that specifically bind to selected antigens (see, *e.g.*, McCafferty *et al.*, *Nature* 348: 552-554 (1990); Marks *et al.*, *Biotechnology* 10: 779-783 (1992)).

Hybridomas can be screened using enzyme-linked immunosorbent assays (ELISA) in order to detect cultures secreting antibodies specific for refolded recombinant CLASP-2. Cultures can also be screened by ELISA to identify those cultures secreting antibodies specific for mammalian-produced CLASP-2. Confirmation of antibody specificity can be obtained by western blot using the same antigens. Subsequent ELISA testing can use recombinant CLASP-2 fragments to identify the specific portion of the CLASP-2 molecule with which a monoclonal antibody binds. Additional testing can be used to identify monoclonal antibodies with desired functional characteristics such as staining of histological sections, immunoprecipitation of CLASP-2, inhibition of CLASP-2 binding or stimulation of CLASP-2 to transmit an intracellular signal. Determination of the monoclonal antibody isotype can be accomplished by ELISA, thus providing additional information concerning purification or function.

Some anti-CLASP-2 monoclonal antibodies of the present invention are humanized, human or chimeric, in order to reduce their potential antigenicity, without reducing their affinity for their target. Humanized antibodies have been described in the art.

See, e.g., Queen, *et al.*, 1989, Proc. Natl Acad. Sci. U.S.A. 86: 10029; U.S. Patent Nos. 5,563,762; 5,693,761; 5,585,089 and 5,530,101. The human antibody sequences used for humanization can be the sequences of naturally occurring human antibodies or can be consensus sequences of several human antibodies. See Kettleborough *et al.*, 1991, *Protein Engineering* 4: 773; Kolbinger *et al.*, 1993, *Protein Engineering* 6: 971. Humanized monoclonal antibodies against CLASP-2 peptides can also be produced using transgenic animals having elements of a human immune system (see, e.g., U.S. Patent Nos. 5,569,825; 5,545,806; 5,693,762; 5,693,761; and 5,7124,350).

In some embodiments, an anti-CLASP-2 polypeptide monoclonal or polyclonal antiserum is produced that is specifically immunoreactive with a particular CLASP-2 polypeptide and is selected to have low cross-reactivity against other molecules (e.g., other CLASP polypeptides) and any such cross-reactivity is removed by immunoabsorbption prior to use in the immunoassay. Methods for screening and characterizing monoclonal antibodies for specificity are well known in the art and are described generally in Harlow and Lane, *supra*. For example, polyclonal antibodies raised to hCLASP-2A, as shown in SEQ ID NO: 1, or splice variants, or immunogenic portions thereof, can be selected to obtain only those polyclonal or monoclonal antibodies that are specifically immunoreactive with the target protein not with other proteins. This selection may be achieved by subtracting out antibodies that cross-react with molecules. A variety of immunoassay formats may be used to select antibodies specifically immunoreactive with a particular protein. For example, solid-phase ELISA immunoassays are routinely used to select antibodies specifically immunoreactive with a protein (see, e.g., Harlow & Lane, *Antibodies, A Laboratory Manual* (1988) for a description of immunoassay formats and conditions that can be used to determine specific immunoreactivity). Typically a specific or selective reaction will be at least twice background signal or noise and more typically more than 10 to 100 times background. Alternatively, antibodies that cross-react with a selected set of polypeptides may be prepared.

Antibody fragments which contain specific binding sites of V can be generated by known techniques. For example, such fragments include, but are not limited to, the F(ab')<sub>2</sub> fragments which can be produced by pepsin digestion of the antibody molecule and the Fab fragments which can be generated by reducing the disulfide bridges of the F(ab')<sub>2</sub> fragments. Alternatively, Fab expression libraries can be constructed (Huse *et al.*, 1989, *Science*, 246: 1275-1281) to allow rapid and easy identification of monoclonal Fab fragments with the desired specificity to CLASP-2.

Anti-CLASP-2 antibodies can also be used to identify, isolate, inhibit or eliminate CLASP-2-expressing cells. In one embodiment, the present invention includes a method of identifying an abnormal T cell profile of an immunocompromised subject relative to the T cell profile of a non-immunocompromised subject. The method includes (i) sorting a sample of peripheral blood mononuclear cells (PBMC) isolated from the immunocompromised subject into sets of T cell types, (ii) determining the ratio of CLASP-2<sup>+</sup> cells relative to the total number of cells (CLASP-2<sup>+</sup>: total) in each set, and identifying an abnormal T cell profile in the immunocompromised subject by comparing the CLASP-2<sup>+</sup>: total ratios of sets from the immunocompromised subject with the CLASP-2<sup>+</sup>: total ratios of analogous sets from a non-immunocompromised subject.

In other embodiments, anti-CLASP-2 antibodies can be used for detection of hCLASP-2 protein in assays such as fluorescent activated cell sorting (FACS), ELISA, fluorescent or electron immunomicroscopy, Western blots, gel shift analyses. CLASP-2 expression in various cells, localization within cells, interactions with other proteins, and differentiation between CLASP-2 isoform expression can be determined by use of the techniques listed herein.

### **5.9. Screening Assays**

The invention provides methods for identifying compounds or agents that modulate (*i.e.*, inhibit or enhance) CLASP-2 expression or activity. CLASP-2 expression or activity modulators are useful for treatment of disorders characterized by (or associated with) aberrant or abnormal CLASP-2 expression or activity. Aberrant expression of CLASP-2 mRNA or protein means expression in lymphocytes (*e.g.*, T lymphocytes or B lymphocytes) or other CLASP-2 expressing cells of at least 2-fold, preferably at least 5-fold greater than expression in control lymphocytes obtained from a healthy subject.

The CLASP-2 expression assays can include the steps of contacting a cell expressing CLASP-2 with a compound or agent and assaying CLASP-2 expression. CLASP-2 polypeptide expression is easily measured by ELISA using anti-CLASP-2 antibodies of the invention. CLASP-2 mRNA expression (including expression of specific species or splice variants of CLASP-2) can be measured by quantitative Northern analysis or quantitative PCR.

CLASP-2 activities include, for example, the CLASP-2 polypeptide binding to PDZ-domain containing molecules and CLASP-2 polypeptide involvement in signal transduction (*e.g.*, leading to T cell activation). Compounds or agents that modulate the

interaction of a CLASP-2 polypeptide and a target molecule, modulate CLASP-2 nucleic acid expression, or modulate CLASP-2 polypeptide activity are all contemplated by the methods of the present invention.

Test compounds include, for example, 1) peptides (*e.g.*, soluble peptides, including Ig-tailed fusion peptides and members of random peptide libraries (see, *e.g.*, Lam, K. S. *et al.*, 1991, Nature 354: 82-84; Houghten, R. *et al.*, 1991, Nature 354: 84-86) and combinatorial chemistry-derived molecular libraries made of D- and/or L-configuration amino acids; 2) phosphopeptides (*e.g.*, members of random and partially degenerate, directed phosphopeptide libraries, see, *e.g.*, Songyang, Z. *et al.*, 1993, Cell 72: 767-778); 3) CLASP-2 antibodies (as described above); 4) small organic and inorganic molecules (*e.g.*, molecules obtained from combinatorial and natural product libraries); 5) antisense RNA and DNA molecules and ribozymes (described above).

The CLASP modulators can be any of a large variety of compounds, both naturally occurring and synthetic, organic and inorganic, and including polymers (*e.g.*, oligopeptides, polypeptides, oligonucleotides, and polynucleotides), small molecules, antibodies, sugars, fatty acids, nucleotides and nucleotide analogs, analogs of naturally occurring structures (*e.g.*, peptide mimetics, nucleic acid analogs, and the like), and numerous other compounds.

In one embodiment, the invention provides assays for screening test compounds which bind to CLASP-2 polypeptides. The assays can be recombinant cell based or cell-free assays. These assays can include the steps of combining a cell expressing a CLASP-2 polypeptide or a binding fragment thereof, and a compound or agent under conditions which allow binding of the compound or agent to the CLASP-2 polypeptide to form a complex. Complex formation can then be determined. The ability of the candidate compound or agent to bind to the CLASP-2 polypeptide or fragment thereof is indicated by the presence of the candidate compound in the complex. Formation of complexes between the CLASP-2 polypeptide and the candidate compound can be quantitated, for example, using standard immunoassays.

In another embodiment, the invention provides screening assays to identify test compounds which modulate the interaction (and most likely CLASP-2 activity as well) between a CLASP-2 polypeptide and a molecule (target molecule with which the CLASP-2 polypeptide normally interacts).

In one embodiment, these CLASP-2 target molecules can be tyrosine kinases (*e.g.*, lyn, lck, fyn, ZAP-70m SyK, and CSK). In another embodiment, these CLASP-2 target

molecules can be tyrosine phosphatases (*e.g.*, EZRIN, SHP-1, SHP-2 and PTP36). In another embodiment, these CLASP-2 target molecules can be adaptor proteins (*e.g.*, NCK, CBL, SHC, LNK, SLP-76, HS1, SIT, VAV, GrB2, and BRDG1). In another embodiment, these CLASP-2 target molecules can be cytoskeletal associated proteins such as ankyrin, spectrin, talin, ezrin, tropomyosin, myosin, plectin, syndecans, paralemmin, Band 3 protein, cytoskeletal protein 4.1, and PTP36. In a further embodiment, CLASP-2 target molecules can be members of the integrin family.

Typically, the assays are recombinant cell based or cell-free assay. These assays can include the steps of combining a cell expressing a CLASP-2 polypeptide or a binding fragment thereof, a CLASP-2 target molecule (*e.g.*, a CLASP-2 ligand) and a test compound, under conditions where but for the presence of the candidate compound, the CLASP-2 polypeptide or biologically active portion thereof binds to the target molecule. Detecting complex formation between the CLASP-2 polypeptide or the binding fragment thereof, the CLASP-2 target molecule and a test compound detecting the formation of a complex which includes the CLASP-2 polypeptide and the target molecule can be accomplished. Detection of complex formation can include direct quantitation of the complex by, for example, measuring inductive effects, such as T cell activation, of the CLASP-2 polypeptide. A significant change, such as a decrease, in the interaction of the CLASP-2 and target molecule (*e.g.*, in the formation of a complex between the CLASP-2 and the target molecule) in the presence of a candidate compound (relative to what is detected in the absence of the candidate compound) is indicative of a modulation of the interaction between the CLASP-2 polypeptide and the target molecule. Modulation of the formation of complexes between the CLASP-2 polypeptide and the target molecule can be quantitated using, for example, an immunoassay. To perform cell free drug screening assays, it is desirable to immobilize either CLASP-2 or its target molecule to facilitate separation of complexes from uncomplexed forms of one or both of the polypeptides, as well as to accommodate automation of the assay. CLASP-2 binding to a target molecule, in the presence and absence of a candidate compound, can be accomplished in any vessel suitable for containing the reactants. Examples of such vessels include microtitre plates, test tubes, and microcentrifuge tubes.

In one embodiment, a fusion polypeptide can be provided which adds a domain that allows the polypeptide to be bound to a matrix. Alternatively, the complexes can be dissociated from the matrix, separated by SDS-PAGE, and the level of CLASP-2-binding



polypeptide found in the bead fraction quantitated from the gel using standard electrophoretic techniques.

Other techniques for immobilizing polypeptides on matrices can also be used in the drug screening assays of the invention. For example, either CLASP-2 or its target molecule can be immobilized utilizing conjugation of biotin and streptavidin. Biotinylated CLASP-2 molecules can be prepared from biotin-NHS (N-hydroxy-succinimide) using techniques well known in the art (*e.g.*, biotinylation kit, Pierce Chemicals, Rockford, Ill.), and immobilized in the wells of streptavidin-coated 96 well plates (Pierce Chemical). Alternatively, antibodies reactive with CLASP-2 but which do not interfere with binding of the polypeptide to its target molecule can be derivatized to the wells of the plate, and CLASP-2 trapped in the wells by antibody conjugation. As described above, preparations of a CLASP-2 -binding polypeptide and a candidate compound are incubated in the CLASP-2 -presenting wells of the plate, and the amount of complex trapped in the well can be quantitated. Methods for detecting such complexes include immunodetection of complexes using antibodies reactive with the CLASP-2 target molecule, or which are reactive with CLASP-2 polypeptide and compete with the target molecule; as well as enzyme-linked assays which rely on detecting an enzymatic activity associated with the target molecule.

One method of drug screening utilizes eukaryotic or prokaryotic host cells which are stably transformed with recombinant DNA molecules expressing the CLASP-2, *e.g.*, the protein having the sequence of SEQ ID NO: 2. Such cells, either in viable or fixed form, can be used for standard ligand/receptor binding assays (see, *e.g.*, Parce *et al.* (1989) Science 246: 243-247; and Owicki *et al.* (1990) Proc. Natl Acad. Sci. U.S.A. 87: 4007-4011, which describe sensitive methods to detect cellular responses. A test compound, often labeled, can be assayed for binding or for competition with another ligand for binding. Viable cells could also be used to screen for the effects of drugs on CLASP-2 mediated functions, *e.g.*, T cell activation, second messenger levels, and others).

In another embodiment, the invention provides a method for identifying a compound (*e.g.*, a screening assay) capable of use in the treatment of a disorder characterized by (or associated with) aberrant or abnormal CLASP-2 nucleic acid expression or CLASP-2 polypeptide activity. This method typically includes the step of assaying the ability of the compound or agent to modulate the expression of the CLASP-2 nucleic acid or the activity of the CLASP-2 polypeptide thereby identifying a compound for treating a disorder characterized by aberrant or abnormal CLASP-2 nucleic acid expression or CLASP-2 polypeptide activity.

Methods for assaying the ability of the compound or agent to modulate the expression of the CLASP-2 nucleic acid or activity of the CLASP-2 polypeptide are typically cell-based assays. For example, cells which are sensitive to ligands which transduce signals via a pathway involving CLASP-2 can be induced to overexpress a CLASP-2 polypeptide in the presence and absence of a candidate compound. Candidate compounds which produce a change in CLASP-2-dependent responses can be identified. In one embodiment, expression of the CLASP-2 nucleic acid or activity of a CLASP-2 polypeptide is modulated in cells and the effects of candidate compounds on the readout of interest (such as T cell activation) are measured. For example, the expression of genes which are up- or down-regulated in response to a CLASP-2-dependent signal cascade can be assayed.

Alternatively, modulators of CLASP-2 expression can be identified in a method where a cell is contacted with a candidate compound and the expression of CLASP-2 mRNA or polypeptide in the cell is determined. The level of expression of CLASP-2 mRNA or polypeptide in the presence of the candidate compound is compared to the level of expression of CLASP-2 mRNA or polypeptide in the absence of the candidate compound. The candidate compound can then be identified as a modulator of CLASP-2 nucleic acid expression based on this comparison. For example, when expression of CLASP-2 mRNA or polypeptide is greater in the presence of the candidate compound than in its absence, the candidate compound is identified as a stimulator of CLASP-2 nucleic acid expression. Alternatively, when CLASP-2 nucleic acid expression is less in the presence of the candidate compound than in its absence, the candidate compound is identified as an inhibitor of CLASP-2 nucleic acid expression. The level of CLASP-2 nucleic acid expression in the cells can be determined by methods described herein for detecting CLASP-2 mRNA or polypeptide.

Modulators of CLASP-2 polypeptide activity and CLASP-2 nucleic acid expression identified according to these drug screening assays can be used to treat, for example, immune disorders. These methods of treatment include the steps of administering the modulators of CLASP-2 polypeptide activity or nucleic acid expression, *e.g.*, in a pharmaceutical composition as described in §5.10.1 below, to a subject in need of such treatment, *e.g.*, a subject with a disorder described herein.

#### **5.10. Therapeutic Administration of CLASP-2 Modulators**

The CLASP-2 protein is expressed in lymphocytes and, as noted *supra*, play a role in regulating T cell and B cell interactions, thus making CLASP-2 activity (*e.g.*, CLASP-

2 binding of regulatory proteins) a target for diagnostic and treatment of immune disorders  
and for modulation of immune function (*e.g.*, T cell activation). Additionally, since CLASP-2  
contains domains capable of transducing an intracellular signal, cell surface CLASP-2 can be  
triggered by an anti- CLASP-2 antibody or soluble CLASP-2 or a fragment thereof in order  
5 to enhance the activation state of a lymphocyte.

#### **5.10.1. Formulation and Route of Administration**

A CLASP-2 polypeptide, a fragment thereof, anti-CLASP-2 antibody,  
CLASP-2 polynucleotide (*e.g.*, antisense or ribozyme), or small molecule agonists or  
antagonists can be administered to a subject *per se* or in the form of a pharmaceutical or  
10 therapeutic composition. Pharmaceutical compositions comprising the proteins of the  
invention can be manufactured by means of conventional mixing, dissolving, granulating,  
dragee-making, levigating, emulsifying, encapsulating, entrapping or lyophilizing processes.  
Pharmaceutical compositions can be formulated in conventional manner using one or more  
physiologically acceptable carriers, diluents, excipients or auxiliaries which facilitate  
15 processing of the protein or active peptides into preparations which can be used  
pharmaceutically. Proper formulation is dependent upon the route of administration chosen.

Currently, there are three major classes of protein-derived cell-penetrating  
peptides that have been used for delivering of proteins into cells and animals (Lindgren, M.;  
*et al.*, 2000, Trends Pharmacol Sci. 21: 99-103). In one embodiment, the CLASP-2 protein  
or fragment (encoding a functional domain of CLASP-2) can be introduced into the cell as a  
fusion protein tied to a transporter protein derived from homeoprotein transcription factors  
such as ANTP. In another embodiment, the CLASP-2 protein or fragment (encoding a  
functional domain of CLASP-2) can be introduced into the cell as a fusion protein tied to  
other transcription factors such as the HIV Tat protein and the herpes simplex virus type 1  
25 (HSV-1) VP22 protein. Members in this family have been widely used in different cellular  
and animal systems (Schwarze, S.; *et al.*; 2000, Trends Pharmacol Sci. 21: 45-48). In another  
embodiment, the CLASP-2 protein or fragment (encoding a functional domain of CLASP-2)  
can be introduced into the cell as a fusion protein tied to peptides derived from signal-  
sequences present in several proteins such as HIV-1 gp41. In other embodiments, there are  
30 several synthetic and/or chemeric cell-penetrating peptides such as transportan and  
Amphiphilic model peptide (Lindgren, M.; *et al.*, 2000, Trends Pharmacol Sci. 21: 99-103)  
that can be used. In another embodiment, the CLASP-2 protein or fragment can be

introduced by using anti-DNA antibodies (see, *e.g.*, Zack, D. J., *et al.*, 1996, J. Immunol. 157: 2082-8

For topical administration the proteins of the invention can be formulated as solutions, gels, ointments, creams, suspensions, and the like. as are well-known in the art.

5           Systemic formulations include those designed for administration by injection, *e.g.*, subcutaneous, intravenous, intramuscular, intrathecal or intraperitoneal injection, as well as those designed for transdermal, transmucosal, oral or pulmonary administration.

For injection, the proteins of the invention can be formulated in aqueous solutions, preferably in physiologically compatible buffers such as Hanks's solution, Ringer's  
10       solution, or physiological saline buffer. The solution can contain formulatory agents such as suspending, stabilizing and/or dispersing agents. Alternatively, the proteins can be in powder form for constitution with a suitable vehicle, *e.g.*, sterile pyrogen-free water, before use.

For transmucosal administration, penetrants appropriate to the barrier to be permeated are used in the formulation. Such penetrants are generally known in the art.

15           For oral administration, a composition can be readily formulated by combining the proteins with pharmaceutically acceptable carriers well known in the art. Such carriers enable the proteins to be formulated as tablets, pills, dragees, capsules, liquids, gels, syrups, slurries, suspensions and the like, for oral ingestion by a patient to be treated. For oral solid formulations such as, for example, powders, capsules and tablets, suitable excipients  
20       include fillers such as sugars, such as lactose, sucrose, mannitol and sorbitol; cellulose preparations such as maize starch, wheat starch, rice starch, potato starch, gelatin, gum tragacanth, methyl cellulose, hydroxypropylmethyl-cellulose, sodium carboxymethylcellulose, and/or polyvinylpyrrolidone (PVP); granulating agents; and binding agents. If desired, disintegrating agents can be added, such as the cross-linked  
25       polyvinylpyrrolidone, agar, or alginic acid or a salt thereof such as sodium alginate.

If desired, solid dosage forms can be sugar-coated or enteric-coated using standard techniques.

For oral liquid preparations such as, for example, suspensions, elixirs and solutions, suitable carriers, excipients or diluents include water, glycols, oils, alcohols, and  
30       the like. Additionally, flavoring agents, preservatives, coloring agents and the like can be added.

For buccal administration, the proteins can take the form of tablets, lozenges, and the like. formulated in conventional manner.

For administration by inhalation, the proteins for use according to the present invention are conveniently delivered in the form of an aerosol spray from pressurized packs or a nebulizer, with the use of a suitable propellant, *e.g.*, dichlorodifluoromethane, trichlorofluoromethane, dichlorotetrafluoroethane, carbon dioxide or other suitable gas. In the case of a pressurized aerosol the dosage unit can be determined by providing a valve to deliver a metered amount. Capsules and cartridges of, *e.g.*, gelatin for use in an inhaler or insufflator can be formulated containing a powder mix of the compound and a suitable powder base such as lactose or starch.

The proteins can also be formulated in rectal or vaginal compositions such as suppositories or retention enemas, *e.g.*, containing conventional suppository bases such as cocoa butter or other glycerides.

In addition to the formulations described previously, the proteins can also be formulated as a depot preparation. Such long acting formulations can be administered by implantation (for example subcutaneously or intramuscularly) or by intramuscular injection. Thus, for example, the proteins can be formulated with suitable polymeric or hydrophobic materials (for example as an emulsion in an acceptable oil) or ion exchange resins, or as sparingly soluble derivatives, for example, as a sparingly soluble salt.

Alternatively, other pharmaceutical delivery systems can be employed. Liposomes and emulsions are well known examples of delivery vehicles that can be used to deliver the proteins or peptides of the invention. Certain organic solvents such as dimethylsulfoxide also can be employed, although usually at the cost of greater toxicity. Additionally, the proteins can be delivered using a sustained-release system, such as semipermeable matrices of solid polymers containing the therapeutic agent. Various sustained-release materials have been established and are well known by those skilled in the art. Sustained-release capsules can, depending on their chemical nature, release the proteins for a few weeks up to over 100 days. Depending on the chemical nature and the biological stability of the therapeutic reagent, additional strategies for protein stabilization can be employed.

As the proteins and peptides of the invention can contain charged side chains or termini, they can be included in any of the above-described formulations as the free acids or bases or as pharmaceutically acceptable salts. Pharmaceutically acceptable salts are those salts which substantially retain the biologic activity of the free bases and which are prepared by reaction with inorganic acids. Pharmaceutical salts tend to be more soluble in aqueous and other protic solvents than are the corresponding free base forms.



### **5.10.2. Effective Dosages**

CLASP-2 polypeptides, CLASP-2 fragments and anti-CLASP-2 antibodies will generally be used in an amount effective to achieve the intended purpose. For use to inhibit an immune response, the proteins of the invention, or pharmaceutical compositions thereof, are administered or applied in a therapeutically effective amount. By therapeutically effective amount is meant an amount effective ameliorate or prevent the symptoms, or prolong the survival of, the patient being treated. Determination of a therapeutically effective amount is well within the capabilities of those skilled in the art, especially in light of the detailed disclosure provided herein.

For systemic administration, a therapeutically effective dose can be estimated initially from *in vitro* assays. For example, a dose can be formulated in animal models to achieve a circulating concentration range that includes the IC<sub>50</sub> as determined in cell culture (*i.e.*, the concentration of test compound that inhibits 50% of CLASP-2 binding interactions). Such information can be used to more accurately determine useful doses in humans.

Initial dosages can also be estimated from *in vivo* data, *e.g.*, animal models, using techniques that are well known in the art. One having ordinary skill in the art could readily optimize administration to humans based on animal data.

Dosage amount and interval can be adjusted individually to provide plasma levels of the proteins which are sufficient to maintain therapeutic effect. Usual patient dosages for administration by injection range from about 0.1 to 5 mg/kg/day, preferably from about 0.5 to 1 mg/kg/day. Therapeutically effective serum levels can be achieved by administering multiple doses each day.

In cases of local administration or selective uptake, the effective local concentration of the proteins can not be related to plasma concentration. One having skill in the art will be able to optimize therapeutically effective local dosages without undue experimentation.

The amount of CLASP-2 administered will, of course, be dependent on the subject being treated, on the subject's weight, the severity of the affliction, the manner of administration and the judgment of the prescribing physician.

The therapy can be repeated intermittently while symptoms detectable or even when they are not detectable. The therapy can be provided alone or in combination with other drugs. In the case of autoimmune disorders, the drugs that can be used in combination

with CLASP-2 or fragments thereof include, but are not limited to, steroid and non-steroid immunosuppressive agents.

### **5.10.3. Toxicity**

Preferably, a therapeutically effective dose of the proteins described herein will provide therapeutic benefit without causing substantial toxicity.

Toxicity of the proteins described herein can be determined by standard pharmaceutical procedures in cell cultures or experimental animals, *e.g.*, by determining the LD<sub>50</sub> (the dose lethal to 50% of the population) or the LD<sub>100</sub> (the dose lethal to 100% of the population). The dose ratio between toxic and therapeutic effect is the therapeutic index.

The data obtained from these cell culture assays and animal studies can be used in formulating a dosage range that is not toxic for use in human. The dosage of the proteins described herein lies preferably within a range of circulating concentrations that include the effective dose with little or no toxicity. The dosage can vary within this range depending upon the dosage form employed and the route of administration utilized. The exact formulation, route of administration and dosage can be chosen by the individual physician in view of the patient's condition. (*See, e.g.*, Fingl *et al.*, 1975, In: The Pharmacological Basis of Therapeutics, Ch.1, p.1).

### **5.11. Binding Assays**

CLASP-2 polypeptides can be used to screen for molecules that bind to CLASP-2 or for molecules to which CLASP-2 binds. The binding of CLASP-2 by the molecule can activate (agonist), increase, inhibit (antagonist), or decrease activity of the CLASP-2 or the molecule bound. Examples of such molecules include antibodies, oligonucleotides, proteins (*e.g.*, receptors), or small molecules. Preferably, the molecule is closely related to the natural ligand of CLASP-2, *e.g.*, a fragment of the ligand, or a natural substrate, a ligand, a structural or functional mimetic. (*See*, Coligan *et al.*, Current Protocols in Immunology 1(2): Chapter 5 (1991).) Similarly, the molecule can be closely-related to the natural receptor to which CLASP-2 binds, or at least, a fragment of the receptor capable of being bound by CLASP-2 (*e.g.*, active site). In either case, the molecule can be rationally designed using known techniques.

Preferably, the screening for these molecules involves producing appropriate cells which express CLASP-2, either as a secreted protein or on the cell membrane. Preferred cells include cells from mammals, yeast, *Drosophila*, or *E. coli*. Cells expressing CLASP-2

(or cell membrane containing the expressed polypeptide) are then preferably contacted with a test compound potentially containing the molecule to observe binding, stimulation, or inhibition of activity of either CLASP-2 or the molecule.

The assay can simply test binding of a candidate compound to CLASP-2, where binding is detected by a label, or in an assay involving competition with a labeled competitor. Further, the assay can test whether the candidate compound results in a signal generated by binding to CLASP-2.

Alternatively, the assay can be carried out using cell-free preparations, polypeptide affixed to a solid support, chemical libraries, or natural product mixtures. The assay can also simply comprise the steps of mixing a candidate compound with a solution containing CLASP-2, measuring CLASP-2 activity or binding, and comparing the CLASP-2 activity or binding to a standard. Preferably, an ELISA assay can measure CLASP-2 level or activity in a sample (*e.g.*, biological sample) using a monoclonal or polyclonal antibody. The antibody can measure CLASP-2 level or activity by either binding, directly or indirectly, to CLASP-2 or by competing with CLASP-2 for a substrate.

In another aspect of the invention, the CLASP-2 polypeptides, or fragments thereof, can be used as “bait proteins” in a two-hybrid assay (see, *e.g.*, U.S. Pat. No. 5,283,317; Zervos *et al.*, 1993, *Cell* 72: 223-232; Madura *et al.*, 1993, *J. Biol. Chem.* 268: 12046-12054; Bartel *et al.*, 1993, *Biotechniques* 14: 920-924; Iwabuchi *et al.*, 1993, *Oncogene* 8: 1693-1696; and Brent WO 94/10300), to identify other proteins, which bind to or interact with CLASP-2 (“CLASP-2-binding proteins” or “CLASP-2-bp”) and modulate CLASP-2 polypeptide activity. Such CLASP-2-binding proteins are also likely to be involved in the propagation of signals by the CLASP-2 polypeptides as, for example, upstream or downstream elements of the CLASP-2 pathway.

All of these above assays can be used as diagnostic or prognostic markers. The molecules discovered using these assays can be used to treat disease or to bring about a particular result in a patient by activating or inhibiting the CLASP-2 molecule. Moreover, the assays can discover agents which can inhibit or enhance the production of CLASP-2 from suitably manipulated cells or tissues.

Therefore, the invention includes a method of identifying compounds or agents that bind to CLASP-2 polypeptides comprising the steps of: (a) contacting a CLASP-2 polypeptide with a compound or agent under conditions which allow binding of the compound to the CLASP-2 polypeptide to form a complex and (b) determining if binding has occurred. Moreover, the invention includes a method of identifying agonists or

antagonists comprising the steps of: (a) incubating a candidate compound with CLASP-2, (b) assaying a biological activity, and (b) determining if a biological activity of CLASP-2 has been altered.

Several methods of automating assays have been developed in recent years so as to permit screening of tens of thousands of compounds in a short period. See, *e.g.*, Fodor *et al.*, 1991, Science 251: 767-773, and other descriptions of chemical diversity libraries, which describe means for testing of binding affinity by a plurality of compounds.

#### **5.12. Other Uses of CLASP-2 Polynucleotides and Polypeptides**

The polynucleotides, polypeptides, polypeptide homologues, modulators, and antibodies described herein can be used in one or more of the following methods: a) drug screening assays; b) diagnostic assays particularly in disease identification, allelic screening and pharmacogenetic testing; and c) pharmacogenomics. A CLASP-2 polypeptide of the invention has one or more of the activities described herein and can thus be used to, for example, modulate an immune response in an immune cell, for example by binding to a CLASP-2 binding partner making it unavailable for binding to the naturally present CLASP-2 polypeptide.

In one embodiment, these CLASP-2 binding partners can be tyrosine kinases (*e.g.*, lyn, lck, fyn, ZAP-70m SyK, and CSK). In another embodiment, these CLASP-2 binding partners can be tyrosine phosphatases (*e.g.*, EZRIN, SHP-1, SHP-2 and PTP36). In another embodiment, these CLASP-2 target molecules can be adaptor proteins (*e.g.*, NCK, CBL, SHC, LNK, SLP-76, HS1, SIT, VAV, GrB2, and BRDG1. In another embodiment, these CLASP-2 binding partners can be cytoskeletal associated proteins such as ankyrin, spectrin, talin, ezrin, tropomyosin, myosin, plectin, syndecans, paralemmin, Band 3 protein, cytoskeletal protein 4.1, and PTP36. In a further embodiment, CLASP-2 binding partners can be members of the integrin family. The isolated nucleic acid molecules of the invention can be used to express CLASP-2 polypeptide (*e.g.*, via a recombinant expression vector in a host cell or in gene therapy applications), to detect CLASP-2 mRNA (*e.g.*, in a biological sample) or a naturally occurring or recombinantly generated genetic mutation in an CLASP-2 gene, and to modulate CLASP-2 activity, as described further below. In addition, the CLASP-2 polypeptides can be used to screen drugs or compounds which modulate CLASP-2 polypeptide activity as well as to treat disorders characterized by insufficient production of CLASP-2 polypeptide or production of CLASP-2 polypeptide forms which have decreased activity compared to wild type CLASP-2. Moreover, the anti-CLASP-2 antibodies of the

invention can be used to detect and isolate an CLASP-2 polypeptide, particularly fragments of CLASP-2 present in a biological sample, and to modulate CLASP-2 polypeptide activity.

### **5.13. Diagnostic Assays**

The invention further provides a method for detecting the presence of CLASP-2, or fragment thereof, in a biological sample. Usually the biological sample contains lymphocytes (*e.g.*, from blood). The method involves contacting the biological sample with a compound or an agent capable of detecting CLASP-2 polypeptide or mRNA such that the presence of CLASP-2 is detected in the biological sample.

A preferred agent for detecting CLASP-2 mRNA is a directly or indirectly labeled nucleic acid probe capable of hybridizing to CLASP-2 mRNA. The nucleic acid probe can be, for example, the full-length CLASP-2 cDNA of SEQ ID NO: 1, or a portion thereof, such as an oligonucleotide of at least 15, 30, 50, 100, 250 or 500 nucleotides in length and sufficient to specifically hybridize under stringent conditions to CLASP-2 mRNA.

A preferred agent for detecting CLASP-2 polypeptide is a directly or indirectly labeled antibody capable of binding to a CLASP-2 polypeptide. Antibodies can be polyclonal, or more preferably, monoclonal. An intact antibody, or a fragment thereof (*e.g.*, Fab or F(ab)<sub>2</sub>) can be used. The term “directly or indirectly”, with regard to the probe or antibody, is intended to encompass direct labeling of the probe or antibody by coupling (*i.e.*, physically linking) a detectable substance to the probe or antibody, as well as indirect labeling of the probe or antibody by reactivity with another reagent that is directly labeled. Examples of indirect labeling include detection of a primary antibody using a fluorescently labeled secondary antibody and end-labeling of a DNA probe with biotin such that it can be detected with fluorescently labeled streptavidin. The detection method of the invention can be used to detect CLASP-2 mRNA or polypeptide in a biological sample in vitro as well as in vivo. For example, in vitro techniques for detection of CLASP-2 mRNA include Northern hybridizations and in situ hybridizations. In vitro techniques for detection of CLASP-2 polypeptide include enzyme linked immunosorbent assays (ELISAs), Western blots, immunoprecipitations and immunofluorescence. Alternatively, CLASP-2 polypeptide can be detected in vivo in a subject by introducing into the subject a labeled anti-CLASP-2 antibody. For example, the antibody can be labeled with a radioactive marker whose presence and location in a subject can be detected by standard imaging techniques. Particularly useful are methods which detect the allelic variant of CLASP-2 expressed in a subject and methods which detect fragments of an CLASP-2 polypeptide in a sample.



The invention also encompasses kits for detecting the presence of CLASP-2 in a biological sample. For example, the kit can comprise a directly or indirectly labeled compound or agent capable of detecting CLASP-2 polypeptide or mRNA in a biological sample; means for determining the amount of CLASP-2 in the sample; and means for comparing the amount of CLASP-2 in the sample with a standard. The compound or agent can be packaged in a suitable container. The kit can further comprise instructions for using the kit to detect CLASP-2 mRNA or polypeptide.

The methods of the invention can also be used to detect naturally occurring genetic mutations in an CLASP-2 gene, thereby determining if a subject with the mutated gene is at risk for a disorder characterized by aberrant or abnormal CLASP-2 nucleic acid expression or CLASP-2 polypeptide activity as described herein. In preferred embodiments, the methods include detecting, in a sample of cells from the subject, the presence or absence of a genetic mutation characterized by at least one of an alteration affecting the integrity of a gene encoding an CLASP-2 polypeptide, or the misexpression of the CLASP-2 gene.

#### **5.14. Biological Activities of CLASP-2**

As described herein, CLASP-2 mediates a variety of cell functions in lymphocytes and other cells. As described herein, a variety of assays are useful for detecting or quantitating CLASP-2 activity, or for identifying agents (including polynucleotides, polypeptides, and antibodies of the invention) that modulate CLASP-2 activity (*i.e.*, biological activity, *e.g.*, binding) or expression. Such agents are useful for treatment of diseases and conditions associated with aberrant CLASP-2 expression or activity. Further, following the guidance provided herein, other CLASP-2-mediated activities can be identified by those of skill using routine assays, such as those described below.

Exemplary assays for CLASP-2 function (or modulation of function) include assays for modulation of an *in vitro* or *in vivo* cell response (*e.g.*, an immune response such as lymphocyte activation, antibody production, inflammation) by detecting a change in an activity (*e.g.*, cytokine production, calcium flux, tyrosine phosphorylation, regulation of early activation markers, cell metabolism, proliferation, and the like, as described below) of cells *in vitro* or *in vivo*. In one embodiment, the cells are lymphocytes.

In one assay, for example, recombinant CLASP-2 protein, peptides, or antibodies corresponding to the CLASP-2 extracellular domain can be mixed directly with T and B cells. Cytokine production by these cells can then be measured and the degree of modulation of the immune response quantitated. Alternatively, antigen-presenting B cells are

mixed with untransfected T cells or T cells that have been transfected with CLASP-2 isoforms. Cytokine production (or calcium flux or other assays in §5.14.3) is measured at the appropriate time to determine the effect of CLASP-2 on such an immune response. In a similar assay, B cells transfected with CLASP-2 constructs are tested for their ability to stimulate a T cell to generate an immune response. Transfected constructs in any of these cases could encode, for example, full or partial length CLASP-2 sequences, or antisense constructs to inhibit translation of endogenous CLASP-2 gene. Any of the examples described herein can be used to stimulate an immune response in the presence or absence of CLASP-2 isoforms or antibodies and assay the resulting effects on immune response by the methods listed in §5.14.3.

#### **5.14.1 Methods for Generating an Immune Response in vitro**

In various assays, an effect of an agent on immune cells is detected using an *in vitro* assay. The degree of an immune response can be measured or quantitated by a number of standard assays including those described below.

In one assay, human peripheral blood mononuclear cells (PBMC), human T cell clones (*e.g.*, Jurkat E6, ATCC TIB-152), EBV-transformed B cell clones (*e.g.*, 9D10, ATCC CRL-8752), antigen-specific T cell clones or lines can be used to examine immune responses in vitro. Activation, enhanced activation or inhibition of activation of these cells or cell lines can be used for the evaluation of potential CLASP therapeutics. Standard methods by which hematopoietic cells are stimulated to undergo activation characteristic of an immune response are, for example:

A) Antigen specific stimulation of immune responses. Either pre-immunized or naïve mouse splenocytes can be generated by standard procedures. In addition, antigen-specific T cell clones and hybridomas (*e.g.*, MBP-specific), and numerous B cell lymphoma cell lines (*e.g.*, CH27), have been previously characterized and are available for the assays discussed below. Antigen specific splenocytes or B-cells can be mixed with specific T-cells in the presence of antigen to generate an immune response. This can be performed in the presence or absence of CLASP-2 to assay whether CLASP-2 modulates the immune response as measured by any of the assays in section 5.14.2.

B) Non-specific T cell activation. The following methods can be used to activate T cells in the absence of antigen: 1) cross-linking T cell receptor (TCR) by addition of antibodies against receptor activation molecules (*e.g.*, TCR, CD3, or CD2) together with antibodies against co-stimulator molecules, for example anti-CD28; 2) activating cell surface

receptors in a non-specific fashion using lectins such as concanavalin A (con A) and phytohemagglutinin (PHA); 3) mimicking cell surface receptor-mediated activation using pharmacological agents that activate protein kinase C (*e.g.*, phorbol esters) and increase cytoplasmic  $\text{Ca}^{2+}$  (*e.g.*, ionomycin).

5 C) Non-specific B cell activation: 1) application of antibodies against cell surface molecules such as IgM, CD20, or CD21. 2) Lipopolysaccharide (LPS), phorbol esters, calcium ionophores and ionomycin can also be used to by-pass receptor triggering.

D) Mixed lymphocyte reaction (MLR). Mix donor PBMC with recipient PBMC to activate lymphocytes by presentation of mismatched tissue antigens, which occurs  
10 in all cases except identical twins.

E) Generation of a specific T cell clone or line that recognizes a particular antigen. A standard approach is to generate tetanus toxin-specific T cells from a donor that has recently been boosted with tetanus toxin. Major histocompatibility complex- (MHC-) matched antigen presenting cells and a source of tetanus toxin are used to maintain antigen  
15 specificity of the cell line or T cell clone (Lanzavecchia, A., *et al.*, 1983, Eur. J. Immun. 13: 733-738).

The anticipated mechanism of action of a CLASP-2 polypeptide or polynucleotide should define the appropriate assay to use to investigate its potential enhancement or inhibition of lymphocyte activation. For example, soluble proteins  
20 containing the CLASP extracellular domain may interfere with the interaction between T cells and antigen presenting cells. Such interaction plays a role in the MLR and in antigen-specific T cell activation, but not in non-specific T or B cell activation. The assays described above have the advantage of several possible detection methods for quantitation.

#### **5.14.2. Methods for Generating an Immune Response in vivo**

25 In various assays, an effect of an agent on immune cells is detected using an *in vivo* assay. The degree of an immune response can be measured or quantitated by a number of standard assays including those described below.

(A) Animal Model for Transplantation Rejection: Ectopic Heart  
Transplantation

30 In one embodiment, a standard animal model for graft versus host rejection is ectopic heart transplantation (Fulmer *et al.*, 1963, Am. J. Anat. 113: 273-281). This method involves using BALB/C mice (either sex, and range from 1–9 months) for transplanting cardiac tissue into a surgically-created pocket on the dorsum for both ears made by slitting

the skin over the auricular artery at the base of the ear. Small curved forceps are forced into the slit, bluntly dissecting between the skin and the cartilage plate. Donor tissue is eased into the base of the pocket near the distal edge of the ear. The auricular artery is used to seal off the opening of the pocket. Within 10 to 14 days pulsatile activity of the transplant should be observed. Gross appearance of the graft, patterns of vacuolar supply to the graft area and pulsatile activity can be easily observed utilizing transilluminated light during the first three weeks post-transplantation. Follow-up can continue for for several months.

(B) Animal model for Autoimmune Disease: Induction of Collagen Induced Arthritis (CIA)

Collagen Induced Arthritis (CIA) is a standard model for studying progression and immune (Courtenay *et al.*, 1980, Nature 283: 666 and Wooley *et al.*, 1981, J. Exp. Med. 154: 688). DBA/a mice can be used as an assay for the in vivo relevance of CLASP-2 in vitro testing potential immune therapeutics. In vivo experiments will be performed to examine the ability of potential therapeutics to prevent CIA. We will use 3-5 mice per group to statistically justify our results.

Once a titer of the potency of collagen type II (CII) is obtained therapeutics can be tested. In one embodiment, three mice will be immunized with three different concentrations of CII 50, 200, and 400 µg per animal (Nabozny *et al.*, 1996, J. Exp. Med., 183: 27-37). To induce CIA, animals can be immunized with an appropriate concentration of CII, determined as described above. One half of a 1:1 ratio of antigen:CFA can be injected at the base of the tail and the remainder equally divided in each hind footpad. Mice can be carefully monitored every day for the onset and progression of CIA throughout the experiment until its termination 12 weeks post-immunization with CII. The pieces of heart transplanted can be approximately 3 X 3 mm in size. The severity of arthritis can be assessed following standard procedures known to one of skill in the art.

**5.14.3 Assay Quantitation**

(A) Tyrosine phosphorylation

Tyrosine phosphorylation of early response proteins such as HS1, PLC-r, ZAP-76, and Vav is an early biochemical event following T cell activation. The tyrosine phosphorylated proteins can be detected by Western blot using antibodies against phosphorylated tyrosine residues. Tyrosine phosphorylation of these early response proteins can be used as a standard assay for T cell activation (J. Biol. Chem., 1997, 272(23): 14562-14570). Any change in the phosphorylation pattern of these or related proteins when immune

responses are generated in the presence of CLASP-2 is indicative of a CLASP-2 modulation of this response.

(B) Intracellular Calcium Flux

The kinetics of intracellular  $\text{Ca}^{2+}$  concentrations are measured over time after stimulation of cells preloaded with a calcium sensitive dye. Upon binding the  $\text{Ca}^{2+}$  indicator dye, Fluor-4 (Molecular Probes), exhibits an increase in fluorescence level using flow cytometry, solution fluorometry, and confocal microscopy. Any change in the level or timing of calcium flux when immune responses are generated in the presence of CLASP-2 is indicative of a CLASP-2 modulation of this response

(C) Regulation of early activation markers

Increased and diminished expression/regulation of early lymphocyte activation marker levels such as CD69, IL-2R, MHC class II, B7, and TCR are commonly measured with fluorescently labeled antibodies using flow cytometry. All antibodies are commercially available. Any change in the expression levels of lymphocyte activation markers when immune responses are generated in the presence of CLASP-2 is indicative of a CLASP-2 modulation of this response.

(D) Increased metabolic activity/acid release

Activation of most known signal transduction pathways trigger increases in acidic metabolites. This reproducible biological event is measured as the rate of acid release using a microphysiometer (Molecular Devices), can be used as an early activation marker when comparing the treatment of cells with potential biological therapeutics (McConnell, H.M. *et al.*, 1992, Science 257: 1906-1912 and McConnell, H.M., 1995, Proc. Natl. Acad. Sci. 92: 2750-2754). Any statistically significant increase or decrease in acid release of CLASP-2-treated sample, as compared to control sample (no treatment), suggest an effect of CLASP-2 on biological function.

(E) Cell proliferation/cell viability assays

(1)  $^3\text{H}$ -thymidine incorporation

Exposure of lymphocytes to antigen or mitogen in vitro induces DNA synthesis and cellular proliferation. The measurement of mitotic activity by  $^3\text{H}$ -thymidine incorporation into newly synthesized DNA is one of the most frequently used assays to quantitative T cell activation. Depending on the cell population and form of stimulation used to activate the T cells, mitotic activity can be measured within 24-72 hrs. in vitro, post  $^3\text{H}$ -thymidine pulse (Mishell, B. B. and S. M. Shiigi, 1980, Selected Methods in Cellular Immunology, W. H. Freeman and Company and Dutton, R. W. and Pearce, J. D., 1962,



Nature 194: 93). Any statistically significant increase or decrease in CPM of CLASP-2-treated sample, as compared to control sample (no treatment), suggest an effect of CLASP-2 on biological function.

(2) MTS [5-(3-carboxymethoxyphenyl)-2-(4,5-dimethylthiazolyl)-3(4-sulfophenyl)tetrazolium, inner salt] is a colorimetric method for determining the number of viable cells in proliferation or cytotoxicity assays (Barltrop, J.A. *et al.*, 1991, Bioorg. & Med. Chem. Lett. 1: 611). 1-5 days after lymphocyte activation, MTS tetrazolium compound, Owen's reagent, is bio-reduced by cells into a colored formazan product that is soluble in tissue culture media. Color intensity is read at 490 nm minus 650 nm using a microplate reader. Any statistically significant increase or decrease in color intensity of CLASP-2-treated sample, as compared to control sample (no treatment), can suggest an effect of CLASP-2 on biological function (Mosmann, T., 1983, J. Immunol. Methods 65: 55 and Barltrop, J.A. *et al.* (1991)).

(3) Bromodeoxyuridine (BrdU), a thymidine analogue, readily incorporates into cells undergoing DNA synthesis. BrdU-pulsed cells are labeled with an enzyme-conjugated anti-BrdU antibody (Gratzner, H.G., 1982, Science 218: 474-475.). A colorimetric, soluble substrate is used to visualize proliferating cells that have incorporated BrdU. Reaction is stopped with sulfuric acid and plate is read at 450 nm using a microplate reader. Any statistically significant increase or decrease in color intensity of CLASP-2-treated sample, as compared to control sample (no treatment), suggest an effect of CLASP-2 on biological function.

#### (F) Apoptosis by Annexin V

Programmed cell death or apoptosis is an early event in a cascade of catabolic reactions leading to cell death. A loss in the integrity of the cell membrane allows for the binding of fluorescently conjugated phosphatidylserine. Stained cells can be measured by fluorescence microscopy and flow cytometry (Vermes, I., 1995, J. Immunol. Methods. 180: 39-52). In one embodiment, any statistically significant increase or decrease in apoptotic cell number of CLASP-2-treated sample, as compared to control sample (no treatment), suggest an effect of CLASP-2 on biological function. For evaluating apoptosis in situ, assays for evaluating cell death in tissue samples can also be used in vivo studies.

#### (G) Quantitation of cytokine production

Cell supernatants harvested after cell stimulation for 16-48 hrs are stored at -80°C until assayed or directly tested for cytokine production. Multiple cytokine assays can be performed on each sample. IL-2, IL-3, IFN- $\gamma$  and other cytokine ELISA Assays are

available for mouse, rat, and human (Endogen, Inc. and BioSource). Cytokine production is measured using a standard two-antibody sandwich ELISA protocol as described by the manufacturer. The presence of horseradish peroxidase is detected with 3, 3', 5' tertamethyl benziidine (TMB) substrate and the reaction is stopped with sulfuric acid. The absorbency at 450 nm is measured using a microplate reader. Any statistically significant increase or decrease in color intensity of CLASP-2-treated sample, as compared to control sample (no treatment), suggest an effect of CLASP-2 on biological function.

(H) NF-AT can be visualized by Immunostaining

T cell activation requires the import of nuclear factor of activated T cells (NFAT) to the nucleus. This translocation of NF-AT can be visualized by immunostaining with anti-NF-AT antibody (Cell 1998, 93: 851-861). Therefore, NF-AT nuclear translocation has been used to assay T cell activation. Similarly, NF-AT/luciferase reporter assays have been used as a standard measurement of T cell activation (MCB 1996, 12: 7151-7160).

(I) ELISA for collagen type II (CII)-specific antibodies (see above for related in vivo assay)

C(II) titers from serum of animals immunized with CLASP-2 can be measured and compared. Both TH1-dependent IgG2a and TH2-dependent IgG1 and IgE CII-specific antibody isotypes will be measured by ELISA. Mouse blood will be obtained by orbital bleed one and two months post-immunization with CII. Samples will be allowed to coagulate and centrifuge to obtain sera, and stored at -80°C until assayed by ELISA. Coat ELISA plates with CII and dilute sera. HRP conjugated goat, isotype specific antibody. Plates are then expose to TMB substrate and read at 450 nm using a microplate reader (Nabozny *et al.*, 1996, J. Exp. Med. 183: 27-37). Any change in the levels of Collagen specific antibodies by colorimetric test when immune responses are generated in the presence of CLASP-2 is indicative of a CLASP-2 modulation of this response.

(J) Antibody Production by ELISPOT Assay

A solid-phase enzyme-linked immunospot (ELISPOT) assay for the quantification of isotype-specific antibody secreting cells (Czerkinsky *et al.*, 1983, J Immunol. Methods. 65: 109-121). Both human and mouse B cells can be tested for isotype and antigen specific antibody production. Although based on a standard ELISA, this technique becomes more sensitive by detecting antibody secretion from single cells. Any change in ELISPOT levels when immune responses are generated in the presence of CLASP-2 is indicative of a CLASP-2 modulation of this response.

(K) Cellular degranulation following IgE cross-linking.

Two cell lines have been obtained from ATCC (MEG01 and HEL-17.92), both of which express the human FCεR1 receptor. FCεR1 is the high affinity receptor for IgE complexes, which when coupled to biotin can be cross-linked with avidin to induce degranulation and histamine release of lymphocytes. Following acylation of the sample, histamine is quantified with an enzyme immunoassay competition assay (Immunotech). Histamine release. A statistically significant increase or decrease in histamine concentration of a CLASP-2 treated sample, as compared to control sample (no treatment), suggest an effect of CLASP-2 on biological function. Any change in frequency of degranulation or histamine levels when immune responses are generated in the presence of CLASP-2 is indicative of a CLASP-2 modulation of this response.

(L) Cellular phenotyping of lymphocytes by flow cytometry and Immunocytochemistry

Determining the tissue distribution of lymphocytes following a pathological disorder can aid in identifying specific organ, tissue and lymphocyte involved in an immune response. Cellular phenotyping of lymphocyte trafficking is generally performed with by flow cytometry and Immunocytochemistry. There are several cluster determination (CD) molecules that are routinely used to identify phenotype, activation kinetics, and regulation events of cells. Any change in levels or distribution of CD molecules when immune responses are generated in the presence of CLASP-2 is indicative of a CLASP-2 modulation of this response.

(M) Structure/Function Assays: Homotypic and/or Heterotypic, Calcium-dependant Cell Adhesion

L929 cells can be transfected with CLASP-2 and Neomycin. G418-resistant clones can be screened for CLASP-expression with anti-CLASP peptide-specific antibodies. These CLASP-expressing clones can then be used to test for homotypic and/or heterotypic calcium dependent cell adhesion using the “cell aggregation assay” described for cadherin molecules (Murphy-Erdosh, C. *et al.*, 1995, J. Cell Biol. 129: 1379-1390). Any change in the levels of cellular aggregation when immune responses are generated in the presence of CLASP-2 is indicative of a CLASP-2 modulation of this response.

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The following cDNA clones described in the Specification and further described in the Examples below have been deposited with the American Type Culture



## **6. EXAMPLES**

### **EXAMPLE 1**

#### **Cloning of CLASP-2**

The cloning of the CLASP gene family has not been a straightforward process.

5 The cloning of each CLASP family member required the use of multiple techniques and resources. CLASP-2 was cloned in the following manner: an expressed sequence tag or EST clone (IMAGE clone 815795, derived from human germinal B cells) was identified based on a BLAST search of human GenBank human EST database using CLASP-1 sequences. IMAGE clone 815795 was sequenced completely. A polynucleotide probe prepared from  
10 815975 sequence was labeled with  $^{32}\text{P}$ -dCTP and used to screen human cDNA libraries including Jurkat (Stratagene) and Ramos B cell cDNA library (James Boulter, UCLA). The screening methods employed were as described in Maniatis *et al.*, 1989, Molecular Cloning A Laboratory Manual, Cold Spring Harbor Laboratory, New York. Several clones were identified and clone C9, with an insert of 3,752 base pairs, was sequenced (ABI dye-sequencing system, PE Applied Biosystems; Perkin-Elmer Corporation, 761 Main Avenue,  
15 Norwalk, CT, U.S.A.). A 5' probe was prepared from C9 sequence and used to rescreen the cDNA libraries. Several clones were isolated, but could not be excised from the phage (Stratagene, CA) without deleting the insert. To circumvent this problem, anchor PCR was performed using M13F primer and CLASP-2 primer (C96AS). The PCR fragment was  
20 cloned using the pGEM-T system (Promega), although initial attempts were unsuccessful. The isolated sequence encompassed additional but incomplete cDNA sequence and was determined to carry at least one mutation that may have allowed it to be propagated in bacteria. Commercial libraries from multiple tissue sources including human placenta, B cell, T cell and peripheral blood were exhaustively screened and re-screened resulting in the  
25 acquisition of only partial cDNAs. Generation of cDNA libraries using oligo dT or CLASP-specific primers also resulted in the acquisition of partial cDNAs. Genomic libraries were screened to obtain a portion of the genomic locus for each of the CLASP genes, and a genomic walk was initiated to obtain 5' exons and extend the cDNA sequence.

To obtain additional 5' CLASP-2 sequence, portions of the cDNA and  
30 genomic sequence from a BAC (Bacterial Artificial Chromosome) genomic library were compared to the NCBI database by BLAST. A genomic clone (Genbank identifier: gi9988160) comprising random, shotgun genomic sequence was identified. Using TFASTX (Pearson and Lipman, PNAS (1988) 85:2444-2448), the amino-terminal sequence of human



CLASP4 was compared to 6 frame translation of gi9988160. Areas of gi9988160 that encoded amino acids with high similarity to CLASP4 amino acid sequence were used to design CLASP-2-specific oligonucleotides for RTPCR (reverse transcriptase polymerase chain reaction according to manufacturers instructions: Reverse transcriptase Gibco/BRL, Taq Polymerase from Sigma). Using oligonucleotides hC2gS5 (nucleotides -66 to -44 of FIG. 11) and C2AS18 (reverse complement of nucleotides 2120 to 2140 of FIG. 11) an RTPCR product of approximately 2.2kb was generated, sequenced (dideoxynucleotide termination sequencing, Beckman Coulter CEQ2000) and shown to be additional human CLASP-2 5' sequence. Further complicating the cloning full-length CLASP cDNA products was the difficulty to clone (and subclone) certain CLASP cDNA products. Standard isolation of some of the CLASP cDNAs from a pure phage population following screening of commercially available cDNA libraries ("ZAP-out" procedure, Stratagene) resulted in no bacterial colonies. Similarly, certain RT-PCR products could not be cloned into standard plasmid vectors. No colonies were isolated by cloning these fragments into vectors lacking promoters, reverse orientations, low copy vectors, or by growth at altered temperatures or levels of antibiotic for plasmid selection (examples: CLASP-7 - HC7gS6 to HC7gAS1 and HC7gS3 to HC7AS14; CLASP-4 - C4P2 to hC4ASTM and C4P2 to HC4AS3'; CLASP-1 - hC1S5' to hC1AS3'Kpn and C1S7 to hC1AS3'Kpn; see Primer Table below). One possibility is that sequences contained within certain regions of CLASP cDNAs are bacteriotoxic and therefore not amenable to cloning. To circumvent these problems direct sequencing of RT-PCR products was performed.

Primer Table

CLASP gene	Sense Primer	Sense sequence	Antisense Primer	Antisense sequence
CLASP-7	HC7gS5	AGGCCTTGTCTCTGTTTACCTG	HC7gAS1	TGTCATGTACTGCACTCGCACAGC
CLASP-7	HC7gS3	ACAGGAACCTGCTGTACGTGTAC	HC7AS14	TCGTGGCTGCACAGGATGCGGGTG
CLASP-4	C4P2	GACCCATTAGGAGGTCTAC	HC4AS3'	CGGGATCCATTGTCACCGTACATCTGC
CLASP-4	C4P2	GACCCATTAGGAGGTCTAC	HC4AS3'	CGGGATCCATTGTCACCGTACATCTGC
CLASP-1	hC1S5'	TATGTCTCAGTCACCTACCTG	HC1AS3'Kpn	CTTGGTACCACTTCAGCACTAGATGAGATG
CLASP-1	C1S7	TCAAGACCAGGGCATGCAAG	HC1AS3'Kpn	CTTGGTACCACTTCAGCACTAGATGAGATG

In-frame stop codons were not present suggesting that the cDNA was not full length. To obtain the 5' terminus of CLASP-2 5' RACE was employed. Antisense oligonucleotides directed against the 5' end of the longest CLASP-2 sequence were generated:

Primers used for human CLASP-2 5' RACE

<u>primer</u>	<u>sequence(5' TO 3')</u>	<u>nucleotide position</u>
HC2RACE1	AAGAGCAGCATCTCCCGTAAACAGTC	-15 to 11
HC2RACE2	TAACAAGCTCTGTGCTTCCTCTTCCG	414 to 443
HC2RACE3	ACCACTTTGTTCGGAAGCTGTCGAAACTC	512 to 540
HC2RACE4	TTTGTACAGCCAGCCATGCTTGGTGATC	634 to 661

RACE was carried out using Generacer kit (Invitrogen) according to manufacturers specifications using polyA selected mRNA from 9D10 B cell tissue culture line. The sequence of the oligonucleotides presented is the reverse complement (*i.e.*,

antisense) of the the CLASP1 cDNA at the indicated position based upon numbering in FIG. 11.

The full length cDNA (presented in FIG. 11) is therefore a compilation of cDNA from cDNA libraries, RTPCR products and 5' RACE products. The sequence of the CLASP-2 cDNA is shown in FIG. 11.

## EXAMPLE 2

### Tissue and Cell Line Expression of the CLASP-2 gene

Multiple Tissue Northern blots were purchased from Clontech; hybridization procedures were followed according to manufacturer's procedures and recommendations.

Human T cell line (Jurkat), human myelomonocyte cells (MV4-11), B cells (9D10), monocytes (THP-1), mouse T cells (3A9), mouse B cells (CH27), human promyelocyte (HL60) and human kidney epithelial cells (293 cell line) were maintained as cultured cell lines. For Multiple Cell Northern, RNA was prepared from cell suspensions using the GIBCO-BRL Trizol system. All steps were performed according to the manufacturer's procedures and recommendations. RNA concentrations were determined by the 260nm/280nm light absorption of the RNA solution. 20 µg RNA was ethanol precipitated and resuspended in formamide/formaldehyde buffer and incubated for 15' at 65°C to eliminate putative secondary structures. RNA samples were run over night on a 1.1% agarose gel containing 1.5% formaldehyde (both gel and running buffer were 20 mM sodium phosphate, pH 7.5). To visualize RNA after gel migration, approx. 0.5 µg ethidium bromide was added to each sample prior to the run together with RNA loading buffer. RNA in the gel was then visualized by 260nm wavelength light. After soaking the gel for 15' in deionized water to reduce the concentration of ethidium bromide in the gel, the RNA was transferred onto Amersham Hybond-N plus membrane by capillary blotting in 20 x SSC buffer for 5 hours. Subsequent to blotting, the membrane was washed in 5 x SSC for 3' and RNA was crosslinked to the membrane by UV light (Stratagene Stratalinker).

A probe which recognizes CLASP-2 isoforms A, B, C, and D (probe HC2.2) was used. Probe HC2.2 encompasses to nucleotides 3920 to 4650 (731 bp long) of CLASP-2A. The HC2.2 probe was prepared using standard labeling kits and desalted using pasteur pipette G-50 Sephadex column in TEN (10 mM Tris-HCl, pH 8.0, 1 mM EDTA, and 100 mM NaCl).

Hybridizations of <sup>32</sup>P dCTP labeled DNA probes to the membrane bound RNAs (multiple tissue and multiple cells) were carried out in CLONTECH EXPRESSHYB

solution, at 68°C and for 1-2 hours. Blots were washed 2 times in 2x SSC 0.1% SDS for 10' each at 50°C and then twice in 0.2 x SSC 0.1% SDS for 10' each at 50°C, followed by a 5' wash in 2xSSC at 50°C. Exposure to KODAK BIOMAX MS film was carried out at minus 80°C using amplifying screens. Typical exposure times were 10 to 36 hours.

### EXAMPLE 3

#### Southern Analysis of CLASP-2

BAC DNA was prepared from E. coli over night cultures using the QIAGEN DNA preparation system. All preps were performed according to the manufacturer's procedures, including the modifications for low copy number DNA constructs. Genomic DNA was prepared from HeLa cells (ATCC #CCL-17) using the methods described by Sambrook, Fritsch and Maniatis (1989); DNA concentrations were determined by the 260nm light absorption of the DNA solution, and aliquots corresponding to 20 microgram (µg) genomic DNA or 2 µg for BAC DNA were used for restriction enzyme digests with Eco RI or Hind III (genomic DNA) or Eco RI and Pst I (BAC DNA). Digests were carried out in 150 microliter volume for 4 hours at 37°C. Digested DNA was ethanol precipitated and the pellet was resuspended in 20 microliter deionized water prior to migration over a 1.2 % agarose gel at 35 V over night. Running buffer was TAE, and the gel contained 0.1 µg ethidium bromide/ml to visualize DNA.

Subsequent to gel separation, DNA was visualized by 260 nm wavelength light. The gel was then washed twice for 20' in denaturing buffer (0.5M NaCl, 0.4 N NaOH) and twice in neutralization buffer (1.5 M NaCl, 0.5 M TRIS pH 8.0). DNA was transferred from the gel onto AMERSHAM HYBOND N membrane by capillary blotting in 20 x SSC for 5 hours. The DNA was crosslinked to the membrane by UV light using a Stratagene Stratalinker.

A probe, HC2.1, which recognizes CLASP-2, was used. Probe HC2.1 encompasses nucleotides 325 to 1126 (802 bp long) of CLASP-2A. The HC2.1 probe was prepared using standard labeling kits and desalted using pasteur pipette G-50 Sephadex column in TEN (10 mM Tris-HCl, pH 8.0, 1 mM EDTA, and 100 mM NaCl). Hybridizations of <sup>32</sup>P dCTP labeled DNA against DNA immobilized onto the membrane were carried out at 65°C overnight in modified CHURCH hybridization solution (7% SDS, 0.5 M sodiumphosphate, 1mM EDTA). Membranes were then exposed to KODAK BIOMAX MS film at minus 80°C. Typical exposure times were 12 hours for genomic DNA southern analysis and 3 hours for BAC DNA Southern analysis.

The genomic DNA southern analysis revealed two fragments (~4.5 kb and 1.85 kb) in the Eco RI digested DNA but three fragments in BACs 4 and 6 DNA. The two major bands are identical in both genomic and BAC DNA (FIG. 7).

#### EXAMPLE 4

##### CLASP-2 Genomic Cloning

Genomic clones of human CLASP-2 were obtained using the Release I high density filters from Genome Systems Inc (cat # FBAC-4434). Two rounds of screening were completed. The first round of screening was carried out using a probe corresponding to nucleotides 3830 to 4558 of the human CLASP-2 cDNA by standard protocols specific by Genome Systems. This screen identified two genomic clones, referred to as AVC BAC4 and 7. A second round of screening using a probe that corresponded to nucleotides 1208 to 1604 of human CLASP-2 cDNA identified clone AVC BAC26. All the clones were partially sequenced to authenticate that they were indeed CLASP-2 genomic clones, to verify exon sequences, and to identify exon/intron boundaries. Oligonucleotides for sequencing the BACs were based upon human CLASP-2 cDNA sequence. Sense and antisense sequencing oligonucleotides were designed along the length of the human CLASP-2 cDNA spaced approximately every 200 nucleotides to ensure a high density of coverage of the corresponding genomic regions. Sequencing reactions with primers and BAC DNA were carried out by standard PCR sequencing using Big Dye termination sequencing mix (ABI). Results from sequence reactions were analyzed using Sequencher software (Genecodes). The results are summarized in FIG. 6.

#### EXAMPLE 5

##### Expression of Recombinant CLASP-2A Polypeptide in Bacterial Cells

Portions of hCLASP-2 were cloned into the GST expression vector pGEX (Pharmacia). These include the region spanning the potential Cadherin processing site through 200 amino acids of the predicted extracellular domain (nucleotide 866 – 1459; GST-EC12; 55 kD fusion) and a portion of the intracellular domain (nucleotide 3230 – 4065; GST-cyto; 57 kD fusion). These regions were amplified using primers at the limits of these sequences on either cDNA clones or cDNA generated from Jurkat or Human Peripheral Blood RNA. Amplified DNA sequences were digested with restriction enzymes for cloning in-frame into GST expression vectors. Fusion proteins were expressed by IPTG induction in DH5 $\alpha$  and purified according to instructions from Pharmacia using glutathione-Sepharose



(Pharmacia). SDS-PAGE gel stained with Coomassie Blue showing induced and uninduced expression of the GST-CLASP-2-cyto construct is shown in FIG. 8. These recombinant proteins were expressed in DH5 $\alpha$  and purified according to instructions from Pharmacia using glutathione-Sepharose. Such recombinant proteins were used to generate antibodies (Josman laboratories) using a AVC Rapid Immunization Protocol.

The full length CLASP can easily be expressed from either the beginning of the hCLASP-2 sequence (in frame with nucleotide 2) or from the first or second methionine (nucleotide 278 or nucleotide 476, underlined in FIG. 1) through to the stop codon (nucleotide 4058). Assuming that the GST moiety has a weight of 26 kD, the total predicted sizes are 180, 168, and 164.5 kD respectively. Alternatively, other bacterial expression systems such as 6CLASP HIS tags, Calmodulin binding protein, maltose binding protein can also be used in a similar manner.

## EXAMPLE 6

### Expression of Recombinant CLASP-2A Polypeptide in Mammalian Cells

#### Example 6A. Secreted fusions

Several portions of the predicted extracellular domain were constructed as hIgG fusions using the CD5gamma-1 expression vector (kindly provided by B. Seed, Harvard University). Polypeptides were cloned into this vector in frame with a CD5 leader sequence that directs the fusion protein into the secretory pathway and in frame with a C-terminal hIgG(Fc) protein. This fusion can be secreted from cell lines such as 293 (Hsieh, J-C., 1999, Nature 398: 431-436). Sense primers with hCLASP-2 sequences beginning at nucleotide 866 and antisense primers at nucleotide 1459 (EC12-IgG), nucleotide 2389 (ECC-IgG) and nucleotide 2857 (ECM-IgG) were used to amplify portions of the extracellular domain for insertion into this vector. Recombinant vectors were purified by Maxiprep (Qiagen) and transfected into 293 EBNA- T cells (kindly provided by B. Seed, Harvard University) by calcium phosphate techniques (Sambrook and Maniatis). After 2-7 days, secreted expression was analyzed by an ELISA against the hIgG fusion using a goat F(ab')<sub>2</sub> anti human IgG(Fc) antibody (Jackson Immunolabs) and Protein-A-HRP (Pierce). Intracellular expression was monitored by immunofluorescence microscopy with a FITC labeled goat anti Human IgG(Fc) antibody (Caltag).

#### Example 6B. Intracellular fusions

Similar methods have been used to construct fusions for expression of full length hCLASP-2 isoforms as well as truncated C-terminal forms in other cell lines such as Jurkat. Recombinant hCLASP-2 fragments were either isolated by digestion of cDNA clones or amplified by primers flanking specific regions (Please provide some specific regions). These can be cloned into expression vectors such as pBJ1-neo (Mark Davis, Stanford University), Peak12 (B. Seed, Harvard University), and pDsRed1-N1 (Clontech). pBJ1-neo and Peak12 allow untagged expression of recombinant proteins and pDsRed1-N1 will allow either untagged or a C-terminal Red fluorescent protein tag. These can be used to generate protein or for expression of various forms for functional analyses.

### **EXAMPLE 7**

#### Antisense Inhibition of CLASP-2 Expression

##### Example 7A. Inhibition of CLASP-2 expression *in vitro*

In this example, inhibition of CLASP-2 expression is examined using an *in vitro* cell-free expression system. To identify the useful antisense oligonucleotides, a series of antisense phosphorothioate oligonucleotides (PS-ODNs), which span portion CLASP-2 sequence, can be systematically assayed for the ability to block CLASP-2 expression *in vitro*.

For inhibition of CLASP-2 expression *in vitro*, a CLASP-2 transcription/expression plasmid can be used according to standard methodology for *in vitro* transcription and translation of sense CLASP-2 RNA. Coupled transcription-translation reactions can be performed with a reticulocyte lysate system (Promega TNTTM) according to standard conditions. Each coupled transcription/translation reaction can include CLASP-2 RNA transcribed from the expression plasmid, and a test antisense polynucleotide at a range of standard test concentrations, as well as the luciferase transcription/translation internal control to normalize each reaction (see, *e.g.*, Sambrook *et al.*, *supra*, Ausubel *et al.*, *supra*). The translation reaction can also be performed with sense CLASP-2 RNA that is synthesized *in vitro* in a separate reaction and then added to the translation reaction. <sup>35</sup>S-Met is included in the reaction to label the translation products. The negative control is performed without added PS-ODN or a sense PS-ODN.

The labeled translation products can be separated by gel electrophoresis and quantitated after exposing the gel to a phosphorimager screen. The amount of CLASP-2 protein expressed in the presence of CLASP-2 specific PS-ODNs can be normalized to the co-expressed luciferase control.

Example 7B. Inhibition of CLASP-2 expression *ex vivo*

A. Reagents

Cells: Jurkat, Clone E6-1 ATCC TIB-152; 9D10 ATCC CRL8752; additional cells from the ATCC or NCI.

Media and solutions: RPMI 1640 medium, BioWhitaker; DMEM/M199 medium, BioWhitaker; EMEM, BioWhitaker; Fetal Bovine Serum, Summit (stored frozen at -20°C, stored thawed at 4°C); Trypsin-EDTA, GIBCO (catalogue #25300-054) (stored frozen at -20°C, stored thawed 4°C; Isoton II (stored at RT); DMSO (stored at RT); oligonucleotides (see Table 1 and FIG. 3, stored in solution at -20°C); PBS (Ca<sup>2+</sup>/Mg<sup>2+</sup> free); TE; 10 mM Tris-HCL, pH 8.0; 1 mM EDTA.

To prepare oligonucleotide stocks: Oligonucleotide nucleotides (PS-ODNs) can dissolved in the appropriate amount of TE to make a concentrated stock solution (1 - 20 mM).

B. Treatment of cells *ex vivo* with antisense CLASP-2 oligonucleotides

Stock cultures of cells in log-phase growth (in T75 flask) can be used. Jurkat, and 9D10 cells are used in this assay. Jurkat and 9D10 are suspension cultures and are passed through dilutions in media. Cell density is measured using a Coulter counter or hemacytometer.

For 6-well dishes,  $1.1 \times 10^5$  cells total per well, 2 ml/well is added. The amount of cells can be scaled up or down proportionally for 12-well, 100 mm, or 150 mm dishes. For example, for 12-well dishes, use  $4.6 \times 10^4$  cells in 2 ml media; for 100 mm dishes use  $6 \times 10^5$  cells in 10 ml media; for 150 mm dishes use  $1.7 \times 10^6$  cells in 35 ml media.

An appropriate number of cells (as described in step 2 above) are collected, centrifuged and resuspended in media containing a range of ODN concentrations. The cells are treated in single, duplicate, or triplicate wells. Control wells are treated with TE or sense ODNs diluted in media.

The suspension cultures are washed and resuspended daily with PS-ODN media.

Suspension cultures are grown for 2-4 days. Cells are washed with PBS and density measured using a Coulter counter or a hemocytometer. If necessary, the cells are replated at  $1.1 \times 10^5$  cells per well, 2 ml media per well, and fed with PS-ODN as described above.

Samples of the cells can also be harvested for analysis to determine the effects of CLASP-2 antisense ODNs. Samples are harvested for RNA and analyzed by either Northern analysis or RT-PCR for the presence of CLASP-2 mRNA. Functional consequences of CLASP-2 antisense ODNs can be analyzed by measuring the ability of Jurkat and 9D10 cells to be activated. Jurkat cells are activated by exposure to anti-CD3 and anti-CD28 crosslinking antibodies, and 9D10 cells are activated by exposure to anti-IgM crosslinking antibody or *P. aeruginosa* lipopolysaccharide. A hallmark of activation, calcium influx, can be measured by flow cytometry. Additionally, ELISA assays can be used to measure Interleukin-2 production from Jurkat cells and secreted IgM can be measured using standard assays from 9D10.

Table 5 below shows exemplary oligonucleotides for this assay:

**Table 5**

Oligo	Sequence 5'-3'	length	notes/comments
1	GAAGGCGATCATCACGT GGCCTTCCATCGC	30-mer	encompasses nucleotides 473-502 and spans the putative initiator methionine (underlined). The function of HC2A, 2B, 2C, and 2E isoforms can be eliminated by this oligonucleotide.
2	GCTTCAAGTAATGACTGG TGCAGAACATCTG	31-mer	Oligonucleotide that should recognize HC2A, 2B, 2D, 2E, and 2F. Encompasses nucleotides 2121-2151. Can be eliminate function of these CLASP-2 isoforms.
3	GCTCCTCCTCAGGCAGGC GCTATGGCTGTGG	34-mer	oligonucleotide specific for HC2C based upon a specific exon found at nucleotide 2927. Can eliminate only HC2D function.
4	GTAGGCCCGGTGCAGCGT GTCATACAGATGG	31-mer	oligonucleotide specific for HC2B, 2C, 2D and 2E based upon specific exon sequence found at nucleotide 3153. Can eliminate function of these CLASP-2 isoforms.
5	GCAATGTCTGAGACTTTC GATCATGAACTATG	32-mer	oligonucleotide specific for HC2A, 2B, 2E, and 2F. Encompasses nucleotides 1987-2018. Can eliminate function of these CLASP-2 isoforms.
6	CAGGAGCTGGTTCTTAAA	18-mer	oligonucleotide specific for HC2A, 2D and 2E. Encompasses nucleotides 2219-2224. Can eliminate function of these CLASP-2 isoforms

Table 5 legend. All nucleotide numeration are relative to Human CLASP-2A (HC2A). See FIG. 2A.

## EXAMPLE 8

### Example 8A. Synthesis of carboxyl-termini PDZ-ligand peptides

The GST-PDZ fusion proteins are made following standard procedures. An exemplary GST-PDZ fusion protein was constructed as follows: A 572 bp fragment encoding

two PDZ domains of the human neDLG gene (Genbank Accession No. U49089.1) was amplified from total Jurkat RNA by RT-PCR according to standard protocols (Sambrook, Fritsch, and Maniatis, 1989, Molecular Cloning – A Laboratory Manual. Cold Spring Harbor Press.) using primers flanked by restriction endonuclease sites for cloning. Fragments were purified by Sephaglas (Pharmacia), digested with the appropriate enzymes, and ligated into the GST expression vector pGEX-3X (Pharmacia) cut with similar enzymes. Recombinant constructs were confirmed by sequencing. Fusion proteins were expressed by IPTG induction in DH5 $\alpha$  and purified using glutathione-Sepharose (Pharmacia) according to instructions from Pharmacia. Excess glutathione was removed using a PD10 desalting column (Pharmacia) and samples were diaconcentrated by placing the protein in dialysis tubing (14,000 MW cutoff) and laying the tubing on polyethylene glycol (3350; Sigma) until volume had been reduced by approximately 50%. Glycerol was then added to 35% final concentration and samples were stored at  $-20^{\circ}\text{C}$ . These recombinant proteins have been used to generate antibodies (Josman laboratories) by standard protocols and for biochemical studies describe herein.

Synthetic peptides corresponding to the carboxyl-terminus of a protein of interest are synthesized by standard resin-based chemistry (*e.g.*, Fmoc), labeled with biotin at the amino-terminus when indicated, and cleaved from the resin using a halide containing acid (*e.g.*, trifluoroacetic acid). The synthetic peptides are then purified by reverse phase high performance liquid chromatography (HPLC) and the identity of the peptides are confirmed by mass spectrometry.

Example 8B. Measurement of CLASP-2 peptide binding to PDZ Domain-containing proteins

The binding of a biotinylated carboxyl-terminal peptide to a GST-PDZ fusion protein is measured as follows:

- (1) GST fusion protein containing one or more PDZ domain(s) is coated onto a protein-binding surface. The protein-binding surface is the surface of a polystyrene plate, which in some cases has been pre-treated by coating with 5  $\mu\text{g/ml}$  of goat-anti-GST polyclonal antibody followed by blocking with excess bovine serum albumin (BSA). The concentration of GST fusion protein used is 5–10  $\mu\text{g/ml}$  and the reaction of the GST fusion protein with the plate is carried out in PBS for 1 – 16 hours at



4°C. If not already blocked, the plate is then blocked with BSA (2% in PBS, 2 hours, 4°C)

(2) The plate is washed with PBS.

(3) The biotinylated peptide (generally 0.2–20 µM) is then added to the plate and allowed to react in PBS/2% BSA buffer with the GST fusion protein for 10 minutes at 4°C followed by 20 minutes at 25°C. In cases where competition between a labeled (biotinylated) and unlabeled (non-biotinylated) peptide is performed, the unlabeled peptide is added immediately prior to adding the labeled peptide.

(4) The plate is washed with PBS.

(1) 0.5 µg/ml streptavidin-HRP conjugate is added to the plate in PBS/2 % BSA buffer and allowed to react for 20 minutes at 4°C.

(6) The plate is washed 5 X with detergent (tween 20) containing solution.

(7) The plate is developed by addition of HRP-substrate solution for 20 minutes at room temperature.

(8) The reaction of the HRP and its substrate is terminated by addition of 1 M sulfuric acid.

(9) The optical density of each well of the plate is read at 450 nm.

In cases where measurement of the apparent affinity of PDZ-ligand interaction is desired, the above procedure is carried out with multiple concentrations of the labeled peptide being used in a single experiment. A plot of binding versus peptide concentration added is then fit to the equation:

$$\text{Binding [peptide]} = \text{Saturation Binding} \times ([\text{peptide}] / ([\text{peptide}] + K_d))$$

where “Binding [peptide]” is the binding of a given concentration of peptide to the GST-PDZ fusion protein minus binding to the GST alone control, “K<sub>d</sub>” is the apparent affinity of the binding reaction, and “Saturation Binding” is computed to allow the best fit of the data to the above equation. The term apparent affinity is used because the reaction may

not reach equilibrium during the duration of the binding reaction in which case the apparent affinity would underestimate the actual affinity (*i.e.*, actual  $K_d < \text{observed } K_d$ ).

## EXAMPLE 9

### Expression of human CLASP -2 in activated T-cells

#### 5            General experimental design

10            The expression profiles of human CLASP-2 in T cells upon T cell activation was determined by Northern analysis. Jurkat E6 lymphoblasts were activated by treatment with anti-CD28, PMA, and Ionomycin. Subsequently, total RNA was extracted from cell aliquots harvested at 0, 1, 2, 4, 8, and 14 hours post activation. The RNA concentration of each preparation was determined by the absorption at 260 nm using a spectrophotometer and concentrations of the different RNA preparations were adjusted such that equal quantities of each RNA preparation could be subjected to Northern analysis. Even gel loading was monitored by ethidium bromide staining of the formaldehyde-agarose gel. Northern membranes were hybridized to radioactively labeled probes corresponding to portions of human CLASP-2 and human beta-actin. Expression levels of CLASP-2 at different time points post T-cell activation are proportional to the radioactive signal generated by hybridization by the CLASP-2 specific radioactively labeled probe that remained bound to the Northern membrane under stringent washing conditions. The entire experiment was done in duplicate.

#### 20            Jurkat E6 cell activation

25            Jurkat E6 cells were maintained and tested in complete IMDM medium supplemented with 2 mM glutamine, 10 mM HEPES, 100 u/mL penicillin, 100 µg/mL streptomycin, 0.1 mM nonessential amino acids, 1 mM sodium pyruvate (Gibco/BRL), 50 µM beta mercaptoethanol (Sigma), and 10% fetal calf serum (Gemini). T cells were activated as described per Fraser et al., using 0.1 g/mL mouse anti-human CD28 monoclonal antibody (PharMingen International catalog number 33741A), 50 ng/mL PMA (Sigma), and 1 µM ionomycin (Calbiochem). Following incubation at 37°C and 5.0% v/v CO<sub>2</sub>, 0.5 x10<sup>6</sup> cells were harvested by centrifugation at 500 x g for 10 minutes (min) at room temperature at 0, 1, 2, 4, 8 and 14 hours post activation and subjected to RNA extraction.

30            For RNA preparation, probe labelling and Northern analysis protocols, see methods and procedures described in Example 2 above. The CLASP-2 specific probe

encompassing nucleotides 5352 to 5922 was generated by PCR from a plasmid containing cloned CLASP-2 cDNA sequences using primers C2S12 and C2AS21.

#### Hybridization, Washing, and Exposure

Blots were washed twice in 2x SSC 0.1% SDS for 10 min each at 60° C and then twice in 0.2x SSC 0.1% SDS for 10 min each at 60° C, followed by a 5' wash in 2xSSC at 60° C. Exposure to KODAK BIOMAX MS film was carried out at minus 80° C using amplifying screens. Typically, exposure times were 10 to 36 hours. Signal intensities on Northern membranes were quantified by the use of a phosphor imager system (STORM, Molecular Dynamics). Signals were counted in the "volume report" mode.

#### Results

CLASP-2 expression levels as determined by Northern analysis (FIG. 14) slightly decrease at 1 hour post activation. The maximum decrease of approximately 36 % is seen at 2 hours post activation. Expression levels augment again at 4 hours post activation but do not attain the level that is seen before activation (0 hours). Intensities of CLASP-2-specific signals on the Northern blot were quantified by phosphor imager analysis. Rectangles were drawn around the areas of CLASP-2-specific signal and total quantity of signal was determined by the "volume report" mode; phosphor imager quantification results of two entirely independent experiments are shown in the diagram (green bars corresponds to Northern blot shown). The above result suggests, that transcriptional control of CLASP-2 expression and T-cell activation are functionally linked to each other.

#### **EXAMPLE 10**

Chromosomal location of CLASP-2 and possible disease associations

CLASP-2 cDNA sequences have been mapped to the genomic clone (GI:9926440, GI:9988160) by use of sequence homology bioinformatics tools BLAST.

Clone (GI:9926440, GI:9988160) has previously been mapped to the chromosomal location 13q12-q13. The literature research reports that the mutations, deletions, rearrangements, disomies and/or breakpoints (in general: chromosomal aberrations) in below listed genes make the genes strong candidates for the onset of the listed diseases/disorders. Because the CLASP-2 gene is localized in the chromosome location 13q12-q13, abnormal CLASP-2 gene regulation or deletion, rearrangement and/or mutations in CLASP-2 locus might be directly or indirectly associated with the onset of the listed diseases. Further, CLASP-2 gene can be used as a genetic probe to detect the abnormality in

regions of these below listed genes and as a diagnostic marker for the related disease/disorders.

<b><i>CANDIDATE GENES</i></b>	<b><i>LOCUS</i></b>	<b><i>RELATED DISEASE/DISORDERS</i></b>
<u>IPF1:Insulin promoter factor1</u>	13q12.1	MODY4: non insulin-dependent juvenile type, Defect in pancreatic islet development and insulin transcription.
BRCA2	13q12.3	BCLL2: B cell lymphoma, deletion encompassing BRCA2 causes B cell lymphoma. BRCA2 is one of the responsible genes for DNA repairing in S phase.
	13q13.1-q14.3	Deletion of these locus causes MDS6: Myelo dysplastic syndrome type 6 including AML.

\*\*\*

The present invention is not to be limited in scope by the exemplified embodiments which are intended as illustrations of single aspects of the invention, and any clones, DNA or amino acid sequences which are functionally equivalent are within the scope of the invention. Indeed, various modifications of the invention in addition to those described herein will become apparent to those skilled in the art from the foregoing description and accompanying drawings. Such modifications are intended to fall within the scope of the appended claims. It is also to be understood that all base pair sizes given for nucleotides are approximate and are used for purposes of description.

All publications and patent documents cited above are hereby incorporated by reference in their entirety for all purposes to the same extent as if each were so individually denoted.

\*\*\*

WHAT IS CLAIMED IS:

1           1.       An isolated CLASP-2 polynucleotide, wherein said polynucleotide is  
2           (a) a polynucleotide that has the sequence of SEQ ID NO: 1, 3, 5 or 9; or  
3           (b) a polynucleotide that hybridizes under stringent hybridization conditions to  
4           (a) and encodes a polypeptide having the sequence of SEQ ID NO: 2, 4, 6 or 10 or an allelic  
5           variant or homologue of a polypeptide having the sequence of SEQ ID NO: 2, 4, 6 or 10; or  
6           (c) a polynucleotide that hybridizes under stringent hybridization conditions to  
7           (a) and encodes a polypeptide with at 25 contiguous residues of the polypeptide of SEQ ID  
8           NO: 2, 4, 6 or 10; or  
9           (d) a polynucleotide that hybridizes under stringent hybridization conditions to  
10          (a) and has at least 12 contiguous bases identical to or exactly complementary to SEQ ID NO:  
11          1, 3, 5 or 9.

1           2.       The polynucleotide of claim 1, wherein said polypeptide specifically  
2           binds to a PDZ domain of PSD95, DLG1 or neDLG.

1           3.       The polynucleotide of claim 2, wherein said polypeptide has a binding  
2           affinity of at least  $10^4 \text{ M}^{-1}$  for binding PSD95, DLG1 or neDLG.

1           4.       The polynucleotide of claim 1 that encodes a polypeptide having the  
2           full-length sequence of SEQ ID NO: 2, 4, 6 or 10.

1           5.       The isolated polynucleotide of claim 1, comprising the cDNA coding  
2           sequence of ATCC Deposit Nos. PTA-1562 and PTA-1563 and PTA-1573.

1           6.       An isolated CLASP-2 polynucleotide comprising a nucleotide  
2           sequence that has at least 90% percent identity to SEQ ID NO: 1, 3, 5 or 9.

1           7.       An isolated polypeptide comprising a nucleotide sequence that has at  
2           least 90% sequence identity to SEQ ID NO: 2, 4, 6 or 10 and is immunologically  
3           crossreactive with SEQ ID NO: 2, 4, 6 or 10 or shares a biological function with native  
4           CLASP-2.

1           8.       A vector comprising the polynucleotide of claim 1.







1 30. A method of detecting a CLASP-2 polypeptide in a sample,  
2 comprising:

3 (a) contacting the sample with a polynucleotide of claim 1 or a polynucleotide  
4 that comprises a sequence of at least 12 nucleotides and is complementary to a contiguous  
5 sequence of the polynucleotide of section (a) of claim 1, and (b) determining whether a  
6 hybridization complex has been formed.

1 31. A method of detecting a CLASP-2 nucleotide in a sample, comprising:  
2 (a) using a polynucleotide that comprises a sequence of at least 12 nucleotides  
3 and is complementary to a contiguous sequence of the polynucleotide of section (a) of claim  
4 1, in an amplification process; and

5 (b) determining whether a specific amplification product has been formed.

1 32. A pharmaceutical composition comprising a polynucleotide of claim 1,  
2 a polypeptide of claim 18, or an antibody of claim 25 and a pharmaceutically acceptable  
3 carrier.

1 33. A method of inhibiting an immune response in a subject comprising:  
2 (a) interfering with the expression of a CLASP-2 gene;  
3 (b) interfering with the ability of a CLASP-2 protein to bind to another cell;  
4 (c) interfering with the ability of a CLASP-2 protein to bind to another protein.

1 34. The method of claim 33, wherein the cell is a T cell or a B cell.

1 35. The method of claim 33 comprising contacting the cell with an  
2 effective amount of a polypeptide which comprises the amino acid sequence of SEQ ID NO:  
3 2, 4, 6 or 10 or a fragment thereof.

1 36. A method of inhibiting an immune response in a subject, comprising  
2 administering to the subject a therapeutically effective amount of an antibody which  
3 specifically binds a polypeptide having the sequence of SEQ ID NO: 2, 4, 6 or 10.

1                    37.     A method of preventing or treating a CLASP-2-mediated disease  
2 comprising administering to a subject in need thereof a therapeutically effective amount of a  
3 pharmaceutical composition of claim 32.

1                    38.     The method claim 37, wherein the CLASP-2-mediated disease is an  
2 autoimmune disease.

1                    39.     A method of treating an autoimmune disease in a subject caused or  
2 exacerbated by increased activity of T<sub>H</sub>1 cells consisting of administering a therapeutically  
3 effective amount of a pharmaceutical composition of claim 32 to the subject.

## CLASP-2 TRANSMEMBRANE PROTEINS

### ABSTRACT OF THE DISCLOSURE

The present invention relates to a cell surface molecule, designated cadherin-like asymmetry protein-2 ("CLASP-2"). In particular, it relates to CLASP-2 polynucleotides, polypeptides,  
5 fusion proteins, and antibodies. The invention also relates to methods of modulating an immune response by interfering with CLASP-2 function.

PA 3103045 v1

10



2 32  
GTT TTA CAC CAT CAC CAA AAC CCA GAA TTT TAT GAT GAG ATT AAA ATA GAG TTG CCC ACT  
val leu his his his gln asn pro glu phe tyr asp glu ile lys ile glu leu pro thr

62 92  
CAG CTG CAT GAA AAG CAC CAC CTG TTG CTC ACA TTC TTC CAT GTC AGC TGT GAC AAC TCA  
gln leu his glu lys his his leu leu leu thr phe phe his val ser cys asp asn ser

122 152  
AGT AAA GGA AGC ACG AAG AAG AGG GAT GTC GTT GAA ACC CAA GTT GGC TAC TCC TGG CTT  
ser lys gly ser thr lys lys arg asp val val glu thr gln val gly tyr ser trp leu

182 212  
CCC CTC CTG AAA GAC GGA AGG GTG GTG ACA AGC GAG CAG CAC ATC CCG GTC TCG GCG AAC  
pro leu leu lys asp gly arg val val thr ser glu gln his ile pro val ser ala asn

242 272  
CTT CCT TCG GGC TAT CTT GGC TAC CAA GAG CTT GGG ATG GGC AGG CAT TAT GGT CCG GAA  
leu pro ser gly tyr leu gly tyr gln glu leu gly met gly arg his tyr gly pro glu

302 332  
ATT AAA TGG GTA GAT GGA GGC AAG CCA CTG CTG AAA ATT TCC ACT CAT CTG GTT TCT ACA  
ile lys trp val asp gly gly lys pro leu leu lys ile ser thr his leu val ser thr

362 392  
GTG TAT ACT CAG GAT CAG CAT TTA CAT AAT TTT TTC CAG TAC TGT CAG AAA ACC GAA TCT  
val tyr thr gln asp gln his leu his asn phe phe gln tyr cys gln lys thr glu ser

422 452  
GGA GCC CAA GCC TTA GGA AAC GAA CTT GTA AAG TAC CTT AAG AGT CTG CAT GCG ATG GAA  
gly ala gln ala leu gly asn glu leu val lys tyr leu lys ser leu his ala met glu

482 512  
GGC CAC GTG ATG ATC GCC TTC TTG CCC ACT ATC CTA AAC CAG CTG TTC CGA GTC CTC ACC  
gly his val met ile ala phe leu pro thr ile leu asn gln leu phe arg val leu thr

542 572  
AGA GCC ACA CAG GAA GAA GTC GCG GTT AAC GTG ACT CGG GTC ATT ATT CAT GTG GTT GCC  
arg ala thr gln glu glu val ala val asn val thr arg val ile ile his val val ala

602 632  
CAG TGC CAT GAG GAA GGA TTG GAG AGC CAC TTG AGG TCA TAT GTT AAG TAC GCG TAT AAG  
gln cys his glu glu gly leu glu ser his leu arg ser tyr val lys tyr ala tyr lys

662 692  
GCT GAG CCA TAT GTT GCC TCT GAA TAC AAG ACA GTG CAT GAA GAA CTG ACC AAA TCC ATG  
ala glu pro tyr val ala ser glu tyr lys thr val his glu glu leu thr lys ser met

FIG. 1

722 752  
ACC ACG ATT CTC AAG CCT TCT GCC GAT TTC CTC ACC AGC AAC AAA CTA CTG AGG TAC TCA  
thr thr ile leu lys pro ser ala asp phe leu thr ser asn lys leu leu arg tyr ser

782 812  
TGG TTT TTC TTT GAT GTA CTG ATC AAA TCT ATG GCT CAG CAT TTG ATA GAG AAC TCC AAA  
trp phe phe phe asp val leu ile lys ser met ala gln his leu ile glu asn ser lys

842 |Cadherin Cleavage| 872  
GTT AAG TTG CTG CGA AAC CAG AGA TTT CCT GCA TCC TAT CAT CAT GCA GCG GAA ACC GTT  
val lys leu leu arg asn gln arg phe pro ala ser tyr his his ala ala glu thr val

902 932  
GTA AAT ATG CTG ATG CCA CAC ATC ACT CAG AAG TTT GGA GAT AAT CCA GAG GCA TCT AAG  
val asn met leu met pro his ile thr gln lys phe gly asp asn pro glu ala ser lys

962 992  
AAC GCG AAT CAT AGC CTT GCT GTC TTC ATC AAG AGA TGT TTC ACC TTC ATG GAC AGG GGC  
asn ala asn his ser leu ala val phe ile lys arg cys phe thr phe met asp arg gly

1022 1052  
TTT GTC TTC AAG CAG ATC AAC AAC TAC ATT AGC TGT TTT GCT CCT GGA GAC CCA AAG ACC  
phe val phe lys gln ile asn asn tyr ile ser cys phe ala pro gly asp pro lys thr

1082 1112  
CTC TTT GAA TAC AAG TTT GAA TTT CTC CGT GTA GTG TGC AAC CAT GAA CAT TAT ATT CCG  
leu phe glu tyr lys phe glu phe leu arg val val cys asn his glu his tyr ile pro

1142 1172  
TTG AAC TTA CCA ATG CCA TTT GGA AAA GGC AGG ATT CAA AGA TAC CAA GAC CTC CAG CTT  
leu asn leu pro met pro phe gly lys gly arg ile gln arg tyr gln asp leu gln leu

1202 1232 |Cadherin EC  
GAC TAC TCA TTA ACA GAT GAG TTC TGC AGA AAC CAC TTC TTG GTG GGA CTG TTA CTG AGG  
asp tyr ser leu thr asp glu phe cys arg asn his phe leu val gly leu leu leu arg

xxx| 1292  
GAG GTG GGG ACA GCC CTC CAG GAG TTC CGG GAG GTC CGT CTG ATC GCC ATC AGT GTG CTC  
glu val gly thr ala leu gln glu phe arg glu val arg leu ile ala ile ser val leu

1322 1352  
AAG AAC CTG CTG ATA AAG CAT TCT TTT GAT GAC AGA TAT GCT TCA AGG AGC CAT CAG GCA  
lys asn leu leu ile lys his ser phe asp asp arg tyr ala ser arg ser his gln ala

1382 1412/471  
AGG ATA GCC ACC CTC TAC CTG CCT CTG TTT GGT CTG CTG ATT GAA AAC GTC CAG CGG ATC  
arg ile ala thr leu tyr leu pro leu phe gly leu leu ile glu asn val gln arg ile

1442 1472  
AAT GTG AGG GAT GTG TCA CCC TTC CCT GTG AAC GCG GGC ATG ACC GTG AAG GAT GAA TCC  
asn val arg asp val ser pro phe pro val asn ala gly met thr val lys asp glu ser

FIG. 1 (cont.)

1502	1532
CTG GCT CTA CCA GCT GTG AAT CCG CTG GTG	ACG CCG CAG AAG GGA AGC ACC CTG GAC AAC
leu ala leu pro ala val asn pro leu val	thr pro gln lys gly ser thr leu asp asn
1562	1592
AGC CTG CAC AAG GAC CTG CTG GGC GCC ATC	TCC GGC ATT GCT TCT CCA TAT ACA ACC TCA
ser leu his lys asp leu leu gly ala ile	ser gly ile ala ser pro tyr thr thr ser
1622	1652
ACT CCA AAC ATC AAC AGT GTG AGA AAT GCT	GAT TCG AGA GGA TCT CTC ATA AGC ACA GAT
thr pro asn ile asn ser val arg asn ala	asp ser arg gly ser leu ile ser thr asp
1682	1712
TCG GGT AAC AGC CTT CCA GAA AGG AAT AGT	GAG AAG AGC AAT TCC CTG GAT AAG CAC CAA
ser gly asn ser leu pro glu arg asn ser	glu lys ser asn ser leu asp lys his gln
1742	1772
CAA AGT AGC ACA TTG GGA AAT TCC GTG GTT	CGC TGT GAT AAA CTT GAC CAG TCT GAG ATT
gln ser ser thr leu gly asn ser val val	arg cys asp lys leu asp gln ser glu ile
1802	1832
AAG AGC CTA CTG ATG TGT TTC CTC TAC ATC	TTA AAG AGC ATG TCT GAT GAT GCT TTG TTT
lys ser leu leu met cys phe leu tyr ile	leu lys ser met ser asp asp ala leu phe
1862	1892
ACA TAT TGG AAC AAG GCT TCA ACA TCT GAA	CTT ATG GAT TTT TTT ACA ATA TCT GAA GTC
thr tyr trp asn lys ala ser thr ser glu	leu met asp phe phe thr ile ser glu val
1922	1952
TGC CTG CAC CAG TTC CAG TAC ATG GGG AAG	CGA TAC ATA GCC AGG AAC CAG GAG GGG TTG
cys leu his gln phe gln tyr met gly lys	arg tyr ile ala arg asn gln glu gly leu
1982	2012
GGA CCC ATA GTT CAT GAT CGA AAG TCT CAG	ACA TTG CCT GTT TCC CGT AAC AGA ACA GGA
gly pro ile val his asp arg lys ser gln	thr leu pro val ser arg asn arg thr gly
2042	2072
ATG ATG CAT GCC AGA TTG CAG CAG CTG GGC	AGC CTG GAT AAC TCT CTC ACT TTT AAC CAC
met met his ala arg leu gln gln leu gly	ser leu asp asn ser leu thr phe asn his
2102	2132
AGC TAT GGC CAC TCG GAC GCA GAT GTT CTG	CAC CAG TCA TTA CTT GAA GCC AAC ATT GCT
ser tyr gly his ser asp ala asp val leu	his gln ser leu leu glu ala asn ile ala
2162	2192
ACT GAG GTT TGC CTG ACA GCT CTG GAC ACG	CTT TCT CTA TTT ACA TTG GCG TTT AAG AAC
thr glu val cys leu thr ala leu asp thr	leu ser leu phe thr leu ala phe lys asn
2222	2252
CAG CTC CTG GCC GAC CAT GGA CAT AAT CCT	CTC ATG AAA AAA GTT TTT GAT GTC TAC CTG
gln leu leu ala asp his gly his asn pro	leu met lys lys val phe asp val tyr leu
2282	2312
TGT TTT CTT CAA AAA CAT CAG TCT GAA ACG	GCT TTA AAA AAT GTC TTC ACT GCC TTA AGG

Fig. 1 (cont.)

cys phe leu gln lys his gln ser glu thr ala leu lys asn val phe thr ala leu arg

2342 2372  
TCC TTA ATT TAT AAG TTT CCC TCA ACA TTC TAT GAA GGG AGA GCG GAC ATG TGT GCG GCT  
ser leu ile tyr lys phe pro ser thr phe tyr glu gly arg ala asp met cys ala ala

2402 2432  
CTG TGT TAC GAG ATT CTC AAG TGC TGT AAC TCC AAG CTG AGC TCC ATC AGG ACG GAG GCC  
leu cys tyr glu ile leu lys cys cys asn ser lys leu ser ser ile arg thr glu ala

2462 2492  
TCC CAG CTG CTC TAC TTC CTG ATG AGG AAC AAC TTT GAT TAC ACT GGA AAG AAG TCC TTT  
ser gln leu leu tyr phe leu met arg asn asn phe asp tyr thr gly lys lys ser phe

2522 2552  
GTC CGG ACA CAT TTG CAA GTC ATC ATA TCT GTC AGC CAG CTG ATA GCA GAC GTT GTT GGC  
val arg thr his leu gln val ile ile ser val ser gln leu ile ala asp val val gly

2582 2612  
ATT GGG GAA ACC AGA TTC CAG CAG TCC CTG TCC ATC ATC AAC AAC TGT GCC AAC AGT GAC  
ile gly glu thr arg phe gln gln ser leu ser ile ile asn asn cys ala asn ser asp

2642 2672  
CGG CTT ATT AAG CAC ACC AGC TTC TCC TCT GAT GTG AAG GAC TTA ACC AAA AGG ATA CGC  
arg leu ile lys his thr ser phe ser ser asp val lys asp leu thr lys arg ile arg

2702 2732  
ACG GTG CTA ATG GCC ACC GCC CAG ATG AAG GAG CAT GAG AAC GAC CCA GAG ATG CTG GTG  
thr val leu met ala thr ala gln met lys glu his glu asn asp pro glu met leu val

2762 2792  
GAC CTC CAG TAC AGC CTG GCC AAA TCC TAT GCC AGC ACG CCC GAG CTC AGG AAG ACG TGG  
asp leu gln tyr ser leu ala lys ser tyr ala ser thr pro glu leu arg lys thr trp

2822 2852 |xxxxxxxxxxxxxxxx Predicted  
CTC GAC AGC ATG GCC AGG ATC CAT GTC AAA AAT GGC GAT CTC TCA GAG GCA GCA ATG TGC  
leu asp ser met ala arg ile his val lys asn gly asp leu ser glu ala ala met cys

Transmembrane Domain xxxxxxxxxxxxxxxxxxxxxxxxxxxxxxxxxxxx|  
TAT GTC CAC GTA ACA GCC CTA GTG GCA GAA TAT CTC ACA CGG AAA GGC GTG TTT AGA CAA  
tyr val his val thr ala leu val ala glu tyr leu thr arg lys gly val phe arg gln

2942 2972  
GGA TGC ACC GCC TTC AGG GTC ATT ACC CCA AAC ATC GAC GAG GAG GCC TCC ATG ATG GAA  
gly cys thr ala phe arg val ile thr pro asn ile asp glu glu ala ser met met glu

3002 3032  
GAC GTG GGG ATG CAG GAT GTC CAT TTC AAC GAG GAT GTG CTG ATG GAG CTC CTT GAG CAG  
asp val gly met gln asp val his phe asn glu asp val leu met glu leu leu glu gln

3062 3092  
TGC GCA GAT GGA CTC TGG AAA GCC GAG CGC TAC GAG CTC ATC GCC GAC ATC TAC AAA CTT

Fig. 1 (cont.)

cys ala asp gly leu trp lys ala glu arg tyr glu leu ile ala asp ile tyr lys leu

3122 3155  
ATC ATC CCC ATT TAT GAG AAG CGG AGG GAT TTC TTT GAA GAT GAA GAT GGA AAG GAG TAT  
ile ile pro ile tyr glu lys arg arg asp phe phe glu asp glu asp gly lys glu tyr

3182 3212  
ATT TAC AAG GAA CCC AAA CTC ACA CCG CTG TCG GAA ATT TCT CAG AGA CTC CTT AAA CTG  
ile tyr lys glu pro lys leu thr pro leu ser glu ile ser gln arg leu leu lys leu

3242 3272  
TAC TCG GAT AAA TTT GGT TCT GAA AAT GTC AAA ATG ATA CAG GAT TCT GGC AAG GTC AAC  
tyr ser asp lys phe gly ser glu asn val lys met ile gln asp ser gly lys val asn

3302 3332  
CCT AAG GAT CTG GAT TCT AAG TAT GCA TAC ATC CAG GTG ACT CAC GTC ATC CCC TTC TTT  
pro lys asp leu asp ser lys tyr ala tyr ile gln val thr his val ile pro phe phe

3362 3392

GAC GAA AAA GAG TTG CAA GAA AGG AAA ACA GAG TTT GAG AGA TCC CAC AAC ATC CGC CGC  
asp glu lys glu leu gln glu arg lys thr glu phe glu arg ser his asn ile arg arg

3422 3452  
TTC ATG TTT GAG ATG CCA TTT ACG CAG ACC GGG AAG AGG CAG GGC GGG GTG GAA GAG CAG  
phe met phe glu met pro phe thr gln thr gly lys arg gln gly gly val glu glu gln

3482 3512

TGC AAA CGG CGC ACC ATC CTG ACA GCC ATA CAC TGC TTC CCT TAT GTG AAG AAG CGC ATC

cys lys arg arg thr ile leu thr ala ile his cys phe pro tyr val lys lys arg ile

```

3542                                     3572 |xxxxxxxxxxxxxxxxxxxxxxxxxxxxxxxx
CCT GTC ATG TAC CAG CAC CAC ACT GAC CTG AAC CCC ATC GAG GTG GCC ATT GAC GAG ATG
pro val met tyr gln his his thr asp leu asn pro ile glu val ala ile asp glu met

```

3602 xxx Coiled-Coil 1 xxxxxxxxxxxxxxxx 3632 xxxx Coiled-Coil 1 xxxxxxxxxxxxxxxx  
AGT AAG AAG GTG GCG GAG CTC CGG CAG CTG TGC TCC TCG GCC GAG GTG GAC ATG ATC AAA  
ser lys lys val ala glu leu arg gln leu cys ser ser ala glu val asp met ile lys

3662 xxxxxxxxxxxxxxxxxxxxxxxxxxxx| 3692  
CTG CAG CTC AAA CTC CAG GGC AGC GTG AGT GTT CAG GTC AAT GCT GGC CCA CTA GCA TAT  
leu gln leu lys leu gln gly ser val ser val gln val asn ala gly pro leu ala tyr

3722											3752									
GCG	CGA	GCT	TTC	TTA	GAT	GAT	ACA	AAC	ACA	AAG	CGA	TAT	CCT	GAC	AAT	AAA	GTG	AAG	CTG	
ala	arg	ala	phe	leu	asp	asp	thr	asn	thr	lys	arg	tyr	pro	asp	asn	lys	val	lys	leu	

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3782                                3812                                |xxxxxxxxxxxxxxxxxxxxxx
CTT AAG GAA GTT TTC AGG CAA TTT GTG GAA GCT TGC GGT CAA GCC TTA GCG GTA AAC GAA
leu lys glu val phe arg gln phe val glu ala cys gly gln ala leu ala val asn glu

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3842 xxx Coiled-Coil 2 xxxxxxxxxxxxxxxx 3872 xx  
CGT CTG ATT AAA GAA GAC CAG CTC GAG TAT CAG GAA GAA ATG AAA GCC AAC TAC AGG GAA

FIG. 1 (cont.)



arg leu ile lys glu asp gln leu glu tyr gln glu glu met lys ala asn tyr arg glu

3902 xxx Coiled-Coil 2 xxxxxxxxxxxxxxxx 3932 xxxxx|  
ATG GCG AAG GAG CTT TCT GAA ATC ATG CAT GAG CAG ATC TGC CCC CTG GAG GAG AAG ACG  
met ala lys glu leu ser glu ile met his glu gln ile cys pro leu glu glu lys thr

3962 3992  
AGC GTC TTA CCG AAT TCC CTT CAC ATC TTC AAC GCC ATC AGT GGG ACT CCA ACA AGC ACA  
ser val leu pro asn ser leu his ile phe asn ala ile ser gly thr pro thr ser thr

4022 |xxxxx PBM xxxxx|  
ATG GTT CAC GGG ATG ACC AGC TCG TCT TCG GTC GTG TGA TTA CAT CTC ATG GCC CGT GTG  
met val his gly met thr ser ser ser ser val val STP

4082 4112  
TGG GGA CTT GCT TTG TCA TTT GCA AAC TCA GGA TGC TTT CCA AAG CCA ATC ACT GGG GAG

4142 4172  
ACC GAG CAC AGG GAG GAC CAA GGG GAA GGG GAG AGA AAG GAA ATA AAG AAC AAC GTT ATT

4202 4232  
TCT TAA CAG ACT TTC TAT AGG AGT TGT AAG AAG GTG CAC ATA TTT TTT TAA ATC TCA CTG

4262 4292  
GCA ATA TTC AAA GTT TTC ATT GTG TCT TAA CAA AGG TGT GGT AGA CAC TCT TGA GCT GGA

4322 4352  
CTT AGA TTT TAT TCT TCC TTG CAG AGT AGT GTT AGA ATA GAT GGC CTA CAG AAA AAA AAG

4382 4412  
GTT CTG GGA TCT ACA TGG CAG GGA GGG CTG CAC TGA CAT TGA TGC CTG GGG GAC CTT TTG

4442 4472  
CCT CGA CTC GTG CCG GAA ATC TGA TCG TAA TCA GGG TAC AGA ACT TAC TAG TTT TGT CTA

4502 4532  
GGA GTA TGT TGT ATG ACT AGG ATT TGT GCT ATT ATC TCA TTC AAC AAC ATA GAG CAA GAA

4562 4592  
TAG TGA GCT AAC TGA GCT AGA CAC TCA ATT AAT CCG CTA CTG GCT TCA AGT CAG AAC TTT

4622 4652  
GTC ATT AAT CAT CGA CTC CGG GAC GGT CAT ATA TGT ATT ACA TTT CTA CAT TTT TAA TAC

4682 4712  
TCA CAT GGG CTT ATG CAT TAA GTT TAA TTG TGA TAA ATT TGT GCT GGT CCA GTA TAT GCA

4742 4772  
ATA CAC TTT AAT GGT TTA TTC TTG TCA TAA AAA TGT GCA ATA TGG AGA TGT ATA CAA GTC

4802  
TTT ACT

Fig. 1 (cont.)

(Nucleotide position for insertions and deletion are found above the Human (h) CLASP-2A line diagram. Numbers are referenced based on hCLASP-2A nucleotide sequence from Figure 1.)

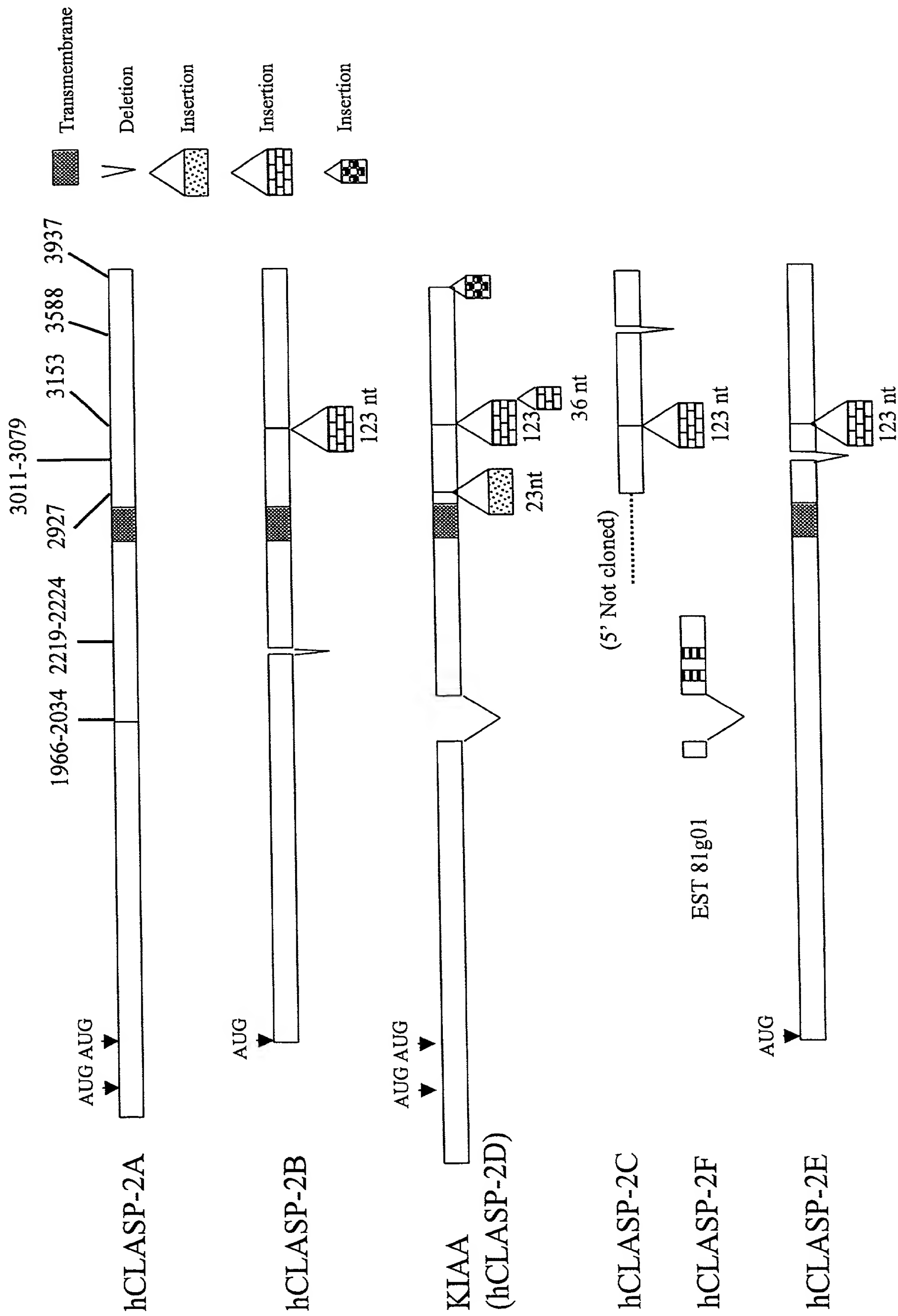


FIG. 2A

2 32  
GTT TTA CAC CAT CAC CAA AAC CCA GAA TTT TAT GAT GAG ATT AAA ATA GAG TTG CCC ACT  
val leu his his his gln asn pro glu phe tyr asp glu ile lys ile glu leu pro thr

62 92  
CAG CTG CAT GAA AAG CAC CAC CTG TTG CTC ACA TTC TTC CAT GTC AGC TGT GAC AAC TCA  
gln leu his glu lys his his leu leu leu thr phe phe his val ser cys asp asn ser

122 152  
AGT AAA GGA AGC ACG AAG AAG AGG GAT GTC GTT GAA ACC CAA GTT GGC TAC TCC TGG CTT  
ser lys gly ser thr lys lys arg asp val val glu thr gln val gly tyr ser trp leu

182 212  
CCC CTC CTG AAA GAC GGA AGG GTG GTG ACA AGC GAG CAG CAC ATC CCG GTC TCG GCG AAC  
pro leu leu lys asp gly arg val val thr ser glu gln his ile pro val ser ala asn

242 272  
CTT CCT TCG GGC TAT CTT GGC TAC CAA GAG CTT GGG ATG GGC AGG CAT TAT GGT CCG GAA  
leu pro ser gly tyr leu gly tyr gln glu leu gly met gly arg his tyr gly pro glu

302 332  
ATT AAA TGG GTA GAT GGA GGC AAG CCA CTG CTG AAA ATT TCC ACT CAT CTG GTT TCT ACA  
ile lys trp val asp gly gly lys pro leu leu lys ile ser thr his leu val ser thr

362 392  
GTG TAT ACT CAG GAT CAG CAT TTA CAT AAT TTT TTC CAG TAC TGT CAG AAA ACC GAA TCT  
val tyr thr gln asp gln his leu his asn phe phe gln tyr cys gln lys thr glu ser

422 452  
GGA GCC CAA GCC TTA GGA AAC GAA CTT GTA AAG TAC CTT AAG AGT CTG CAT GCG ATG GAA  
gly ala gln ala leu gly asn glu leu val lys tyr leu lys ser leu his ala met glu

482 512  
GGC CAC GTG ATG ATC GCC TTC TTG CCC ACT ATC CTA AAC CAG CTG TTC CGA GTC CTC ACC  
gly his val met ile ala phe leu pro thr ile leu asn gln leu phe arg val leu thr

542 572  
AGA GCC ACA CAG GAA GAA GTC GCG GTT AAC GTG ACT CGG GTC ATT ATT CAT GTG GTT GCC  
arg ala thr gln glu glu val ala val asn val thr arg val ile ile his val val ala

602 632  
CAG TGC CAT GAG GAA GGA TTG GAG AGC CAC TTG AGG TCA TAT GTT AAG TAC GCG TAT AAG  
gln cys his glu glu gly leu glu ser his leu arg ser tyr val lys tyr ala tyr lys

662 692  
GCT GAG CCA TAT GTT GCC TCT GAA TAC AAG ACA GTG CAT GAA GAA CTG ACC AAA TCC ATG  
ala glu pro tyr val ala ser glu tyr lys thr val his glu glu leu thr lys ser met

FIG. 2B

722 752  
ACC ACG ATT CTC AAG CCT TCT GCC GAT TTC CTC ACC AGC AAC AAA CTA CTG AGG TAC TCA  
thr thr ile leu lys pro ser ala asp phe leu thr ser asn lys leu leu arg tyr ser

782 812  
TGG TTT TTC TTT GAT GTA CTG ATC AAA TCT ATG GCT CAG CAT TTG ATA GAG AAC TCC AAA  
trp phe phe phe asp val leu ile lys ser met ala gln his leu ile glu asn ser lys

842 |Cadherin Cleavage| 872  
GTT AAG TTG CTG CGA AAC CAG AGA TTT CCT GCA TCC TAT CAT CAT GCA GCG GAA ACC GTT  
val lys leu leu arg asn gln arg phe pro ala ser tyr his his ala ala glu thr val

902 932  
GTA AAT ATG CTG ATG CCA CAC ATC ACT CAG AAG TTT GGA GAT AAT CCA GAG GCA TCT AAG  
val asn met leu met pro his ile thr gln lys phe gly asp asn pro glu ala ser lys

962 992  
AAC GCG AAT CAT AGC CTT GCT GTC TTC ATC AAG AGA TGT TTC ACC TTC ATG GAC AGG GGC  
asn ala asn his ser leu ala val phe ile lys arg cys phe thr phe met asp arg gly

1022 1052  
TTT GTC TTC AAG CAG ATC AAC AAC TAC ATT AGC TGT TTT GCT CCT GGA GAC CCA AAG ACC  
phe val phe lys gln ile asn asn tyr ile ser cys phe ala pro gly asp pro lys thr

1082 1112  
CTC TTT GAA TAC AAG TTT GAA TTT CTC CGT GTA GTG TGC AAC CAT GAA CAT TAT ATT CCG  
leu phe glu tyr lys phe glu phe leu arg val val cys asn his glu his tyr ile pro

1142 1172  
TTG AAC TTA CCA ATG CCA TTT GGA AAA GGC AGG ATT CAA AGA TAC CAA GAC CTC CAG CTT  
leu asn leu pro met pro phe gly lys gly arg ile gln arg tyr gln asp leu gln leu

1202 1232 |Cadherin EC  
GAC TAC TCA TTA ACA GAT GAG TTC TGC AGA AAC CAC TTC TTG GTG GGA CTG TTA CTG AGG  
asp tyr ser leu thr asp glu phe cys arg asn his phe leu val gly leu leu leu arg

xxx1 1292  
GAG GTG GGG ACA GCC CTC CAG GAG TTC CGG GAG GTC CGT CTG ATC GCC ATC AGT GTG CTC  
glu val gly thr ala leu gln glu phe arg glu val arg leu ile ala ile ser val leu

1322 1352  
AAG AAC CTG CTG ATA AAG CAT TCT TTT GAT GAC AGA TAT GCT TCA AGG AGC CAT CAG GCA  
lys asn leu leu ile lys his ser phe asp asp arg tyr ala ser arg ser his gln ala

1382 1412  
AGG ATA GCC ACC CTC TAC CTG CCT CTG TTT GGT CTG CTG ATT GAA AAC GTC CAG CGG ATC  
arg ile ala thr leu tyr leu pro leu phe gly leu leu ile glu asn val gln arg ile

1442 1472  
AAT GTG AGG GAT GTG TCA CCC TTC CCT GTG AAC GCG GGC ATG ACC GTG AAG GAT GAA TCC  
asn val arg asp val ser pro phe pro val asn ala gly met thr val lys asp glu ser

FIG. 2B (cont.)

1502	1532	
CTG GCT CTA CCA GCT GTG AAT CCG CTG GTG	ACG CCG CAG AAG GGA AGC ACC CTG GAC AAC	
leu ala leu pro ala val asn pro leu val	thr pro gln lys gly ser thr leu asp asn	
1562	1592	
AGC CTG CAC AAG GAC CTG CTG GGC GCC ATC	TCC GGC ATT GCT TCT CCA TAT ACA ACC TCA	
ser leu his lys asp leu leu gly ala ile	ser gly ile ala ser pro tyr thr thr ser	
1622	1652	
ACT CCA AAC ATC AAC AGT GTG AGA AAT GCT	GAT TCG AGA GGA TCT CTC ATA AGC ACA GAT	
thr pro asn ile asn ser val arg asn ala	asp ser arg gly ser leu ile ser thr asp	
1682	1712	
TCG GGT AAC AGC CTT CCA GAA AGG AAT AGT	GAG AAG AGC AAT TCC CTG GAT AAG CAC CAA	
ser gly asn ser leu pro glu arg asn ser	glu lys ser asn ser leu asp lys his gln	
1742	1772	
CAA AGT AGC ACA TTG GGA AAT TCC GTG GTT	CGC TGT GAT AAA CTT GAC CAG TCT GAG ATT	
gln ser ser thr leu gly asn ser val val	arg cys asp lys leu asp gln ser glu ile	
1802	1832	
AAG AGC CTA CTG ATG TGT TTC CTC TAC ATC	TTA AAG AGC ATG TCT GAT GAT GCT TTG TTT	
lys ser leu leu met cys phe leu tyr ile	leu lys ser met ser asp asp ala leu phe	
1862	1892	
ACA TAT TGG AAC AAG GCT TCA ACA TCT GAA	CTT ATG GAT TTT TTT ACA ATA TCT GAA GTC	
thr tyr trp asn lys ala ser thr ser glu	leu met asp phe phe thr ile ser glu val	
1922	1952	xxxxxxxxxxxxxxxxxxxxxx
TGC CTG CAC CAG TTC CAG TAC ATG GGG AAG	CGA TAC ATA GCC AGG AAC CAG GAG GGG TTG	
cys leu his gln phe gln tyr met gly lys	arg tyr ile ala arg asn gln glu gly leu	
1982	xxxxxxxxxx deleted in CLASP-2D(KIAA1058) xxxxxxxxxxxxxxxxxxxxxxxxx	
GGA CCC ATA GTT CAT GAT CGA AAG TCT CAG	ACA TTG CCT GTT TCC CGT AAC AGA ACA GGA	
gly pro ile val his asp arg lys ser gln	thr leu pro val ser arg asn arg thr gly	
2042	2072	
ATG ATG CAT GCC AGA TTG CAG CAG CTG GGC	AGC CTG GAT AAC TCT CTC ACT TTT AAC CAC	
met met his ala arg leu gln gln leu gly	ser leu asp asn ser leu thr phe asn his	
2102	2132	
AGC TAT GGC CAC TCG GAC GCA GAT GTT CTG	CAC CAG TCA TTA CTT GAA GCC AAC ATT GCT	
ser tyr gly his ser asp ala asp val leu	his gln ser leu leu glu ala asn ile ala	
		Deleted
2162	2192	lxxx
ACT GAG GTT TGC CTG ACA GCT CTG GAC ACG	CTT TCT CTA TTT ACA TTG GCG TTT AAG AAC	
thr glu val cys leu thr ala leu asp thr	leu ser leu phe thr leu ala phe lys asn	
in HC2B		
xxxl	2252	
CAG CTC CTG GCC GAC CAT GGA CAT AAT CCT	CTC ATG AAA AAA GTT TTT GAT GTC TAC CTG	
gln leu leu ala asp his gly his asn pro	leu met lys lys val phe asp val tyr leu	



2282	2312	
TGT TTT CTT CAA AAA CAT CAG TCT GAA ACG GCT TTA AAA AAT GTC TTC ACT GCC TTA AGG		
cys phe leu gln lys his gln ser glu thr ala leu lys asn val phe thr ala leu arg		
2342	2372	
TCC TTA ATT TAT AAG TTT CCC TCA ACA TTC TAT GAA GGG AGA GCG GAC ATG TGT GCG GCT		
ser leu ile tyr lys phe pro ser thr phe tyr glu gly arg ala asp met cys ala ala		
2402	2432	
CTG TGT TAC GAG ATT CTC AAG TGC TGT AAC TCC AAG CTG AGC TCC ATC AGG ACG GAG GCC		
leu cys tyr glu ile leu lys cys cys asn ser lys leu ser ser ile arg thr glu ala		
2462	2492	
TCC CAG CTG CTC TAC TTC CTG ATG AGG AAC AAC TTT GAT TAC ACT GGA AAG AAG TCC TTT		
ser gln leu leu tyr phe leu met arg asn asn phe asp tyr thr gly lys lys ser phe		
2522	2552	
GTC CGG ACA CAT TTG CAA GTC ATC ATA TCT GTC AGC CAG CTG ATA GCA GAC GTT GTT GGC		
val arg thr his leu gln val ile ile ser val ser gln leu ile ala asp val val gly		
2582	2612	
ATT GGG GAA ACC AGA TTC CAG CAG TCC CTG TCC ATC ATC AAC AAC TGT GCC AAC AGT GAC		
ile gly glu thr arg phe gln gln ser leu ser ile ile asn asn cys ala asn ser asp		
2642	2672	
GGG CTT ATT AAG CAC ACC AGC TTC TCC TCT GAT GTG AAG GAC TTA ACC AAA AGG ATA CGC		
arg leu ile lys his thr ser phe ser ser asp val lys asp leu thr lys arg ile arg		
2702	2732	
ACG GTG CTA ATG GCC ACC GCC CAG ATG AAG GAG CAT GAG AAC GAC CCA GAG ATG CTG GTG		
thr val leu met ala thr ala gln met lys glu his glu asn asp pro glu met leu val		
2762	2792	
GAC CTC CAG TAC AGC CTG GCC AAA TCC TAT GCC AGC ACG CCC GAG CTC AGG AAG ACG TGG		
asp leu gln tyr ser leu ala lys ser tyr ala ser thr pro glu leu arg lys thr trp		
2822	2852	1xxxxxxxxxxxxxxxx Predicted
CTC GAC AGC ATG GCC AGG ATC CAT GTC AAA AAT GGC GAT CTC TCA GAG GCA GCA ATG TGC		
leu asp ser met ala arg ile his val lys asn gly asp leu ser glu ala ala met cys		

[Additional and differential exon usage found at position 2927 consisting of 69 nucleotides. This entire sequence is found in Human CLASP-2D (KIAA1058) and not other isoforms of CLASP-2. It has a sequence of:  
AAGCAGTCCAGTGGGAGCCGCCCTTCTCCCCACAGCCATAGCGCCTGCCTGAGGAGGAGCCGGGGAG]

Transmembrane Domain xxxxxxxxxxxxxxxxxxxxxxxxxxxxxxxx|

TAT GTC CAC GTA ACA GCC CTA GTG GCA GAA TAT CTC ACA CGG AAA GGC GTG TTT AGA CAA  
tyr val his val thr ala leu val ala glu tyr leu thr arg lys gly val phe arg gln

2942	2972
GGA TGC ACC GCC TTC AGG GTC ATT ACC CCA AAC ATC GAC GAG GAG GCC TCC ATG ATG GAA	
gly cys thr ala phe arg val ile thr pro asn ile asp glu glu ala ser met met glu	

Fig. 2B (cont.)



3602 3632  
AGT AAG AAG GTG GCG GAG CTC CGG CAG CTG TGC TCC TCG GCC GAG GTG GAC ATG ATC AAA  
ser lys lys val ala glu leu arg gln leu cys ser ser ala glu val asp met ile lys

3662 3692  
CTG CAG CTC AAA CTC CAG GGC AGC GTG AGT GTT CAG GTC AAT GCT GGC CCA CTA GCA TAT  
leu gln leu lys leu gln gly ser val ser val gln val asn ala gly pro leu ala tyr

3722 3752  
GCG CGA GCT TTC TTA GAT GAT ACA AAC ACA AAG CGA TAT CCT GAC AAT AAA GTG AAG CTG  
ala arg ala phe leu asp asp thr asn thr lys arg tyr pro asp asn lys val lys leu

3782 3812  
CTT AAG GAA GTT TTC AGG CAA TTT GTG GAA GCT TGC GGT CAA GCC TTA GCG GTA AAC GAA  
leu lys glu val phe arg gln phe val glu ala cys gly gln ala leu ala val asn glu

3842 3872  
CGT CTG ATT AAA GAA GAC CAG CTC GAG TAT CAG GAA GAA ATG AAA GCC AAC TAC AGG GAA  
arg leu ile lys glu asp gln leu glu tyr gln glu glu met lys ala asn tyr arg glu

Insertion of 8 nucleotides found only in Human CLASP-2D with sequence: CTGGGATG

3902 3932  
ATG GCG AAG GAG CTT TCT GAA ATC ATG CAT GAG CAG ATC TGC CCC CTG GAG GAG AAG ACG  
met ala lys glu leu ser glu ile met his glu gln ile cys pro leu glu glu lys thr

3962 3992  
AGC GTC TTA CCG AAT TCC CTT CAC ATC TTC AAC GCC ATC AGT GGG ACT CCA ACA AGC ACA  
ser val leu pro asn ser leu his ile phe asn ala ile ser gly thr pro thr ser thr

4022 1xxxx PBM xxxxi  
ATG GTT CAC GGG ATG ACC AGC TCG TCT TCG GTC GTG TGA TTA CAT CTC ATG GCC CGT GTG  
met val his gly met thr ser ser ser ser val val STP

4082 4112  
TGG GGA CTT GCT TTG TCA TTT GCA AAC TCA GGA TGC TTT CCA AAG CCA ATC ACT GGG GAG

4142 4172  
ACC GAG CAC AGG GAG GAC CAA GGG GAA GGG GAG AGA AAG GAA ATA AAG AAC AAC GTT ATT

4202 4232  
TCT TAA CAG ACT TTC TAT AGG AGT TGT AAG AAG GTG CAC ATA TTT TTT TAA ATC TCA CTG

4262 4292  
GCA ATA TTC AAA GTT TTC ATT GTG TCT TAA CAA AGG TGT GGT AGA CAC TCT TGA GCT GGA

4322 4352  
CTT AGA TTT TAT TCT TCC TTG CAG AGT AGT GTT AGA ATA GAT GGC CTA CAG AAA AAA AAG

4382 4412  
GTT CTG GGA TCT ACA TGG CAG GGA GGG CTG CAC TGA CAT TGA TGC CTG GGG GAC CTT TTG

4442 4472

Fig. 2B (cont.)

CCT CGA CTC GTG CCG GAA ATC TGA TCG TAA TCA GGG TAC AGA ACT TAC TAG TTT TGT CTA

4502

4532

GGA GTA TGT TGT ATG ACT AGG ATT TGT GCT ATT ATC TCA TTC AAC AAC ATA GAG CAA GAA

4562

4592

TAG TGA GCT AAC TGA GCT AGA CAC TCA ATT AAT CCG CTA CTG GCT TCA AGT CAG AAC TTT

4622

4652

GTC ATT AAT CAT CGA CTC CCG GAC GGT CAT ATA TGT ATT ACA TTT CTA CAT TTT TAA TAC

4682

4712

TCA CAT GGG CTT ATG CAT TAA GTT TAA TTG TGA TAA ATT TGT GCT GGT CCA GTA TAT GCA

4742

4772

ATA CAC TTT AAT GGT TTA TTC TTG TCA TAA AAA TGT GCA ATA TGG AGA TGT ATA CAA GTC

4802

TTT ACT

05687837 101300

Fig. 2B(cont.)

HC2A	-----
HC2-80	-----
HC2B	-----
HC2C	-----
HC2D-KIAA1058	GCATCTGGAAATCTTGACAAAAATGCCAGATTTTCTGCCATCTACAGGCAAGACAGCAAT
HC2E	-----
HC2F	-----

HC2A	-----
HC2-80	-----
HC2B	-----
HC2C	-----
HC2D-KIAA1058	AAGCTATCCAATGATGACATGCTCAAGTTACTTGCAGACTTTCGGAAACCTGAGAAGATG
HC2E	-----
HC2F	-----

HC2A	-----
HC2-80	-----
HC2B	-----
HC2C	-----
HC2D-KIAA1058	GCTAAGCTCCCAGTGATTTTAGGCAATCTAGACATTACAATTGATAATGTTTCCTCAGAC
HC2E	-----
HC2F	-----

HC2A	-----
HC2-80	-----
HC2B	-----
HC2C	-----
HC2D-KIAA1058	TTCCCTAATTATGTTAATTCATCATACTTCCCACAAAACAATTTGAAACCTGCAGTAAA
HC2E	-----
HC2F	-----

HC2A	-----
HC2-80	-----
HC2B	-----
HC2C	-----
HC2D-KIAA1058	ACTCCCATCACGTTTGAAGTGGAGGAATTTGTGCCCTGCATACCAAACACACTCAGCCT
HC2E	-----
HC2F	-----

FIG. 3A



HC2A	-----
HC2-80	-----
HC2B	-----
HC2C	-----
HC2D-KIAA1058	TACACCATCTACACCAATCACCTTTACGTTTATCCTAAGTACTTGAAATACGACAGTCAG
HC2E	-----
HC2F	-----
HC2A	-----
HC2-80	-----
HC2B	-----
HC2C	-----
HC2D-KIAA1058	AAGTCTTTTGCCAAGGCTAGAAATATTGCGATTTGCATTGAATTCAAAGATTCAGATGAG
HC2E	-----
HC2F	-----
HC2A	-----
HC2-80	-----
HC2B	-----
HC2C	-----
HC2D-KIAA1058	GAAGACTCTCAGCCCCTTAAGTGCATTTATGGCAGACCTGGTGGGCCAGTTTTTACAAGA
HC2E	-----
HC2F	-----
HC2A	-----AGTTTTACACCATCACCAAAACCCAGAATTTTATGATGAGATTAAA
HC2-80	-----
HC2B	-----
HC2C	-----
HC2D-KIAA1058	AGCGCCTTTGCTGCAGTTTTACACCATCACCAAAACCCAGAATTTTATGATGAGATTAAA
HC2E	-----
HC2F	-----
HC2A	ATAGAGTTGCCCCTCAGCTGCATGAAAAGCACCACTGTTGCTCACATTCTTCCATGTC
HC2-80	-----
HC2B	-----
HC2C	-----
HC2D-KIAA1058	ATAGAGTTGCCCCTCAGCTGCATGAAAAGCACCACTGTTGCTCACATTCTTCCATGTC
HC2E	-----
HC2F	-----
HC2A	AGCTGTGACAACTCAAGTAAAGGAAGCACGAAGAAGAGGGATGTCGTTGAAACCCAAGTT
HC2-80	-----
HC2B	-----
HC2C	-----
HC2D-KIAA1058	AGCTGTGACAACTCAAGTAAAGGAAGCACGAAGAAGAGGGATGTCGTTGAAACCCAAGTT
HC2E	-----
HC2F	-----

FIG. 3A (cont.)

HC2A	GGCTACTCCTGGCTTCCCCTCCTGAAAGACGGAAGGGTGGTGACAAGCGAGCAGCACATC
HC2-80	-----
HC2B	-----
HC2C	-----
HC2D-KIAA1058	GGCTACTCCTGGCTTCCCCTCCTGAAAGACGGAAGGGTGGTGACAAGCGAGCAGCACATC
HC2E	-----
HC2F	-----

HC2A	CCGGTCTCGGCGAACCTTCCTTCGGGCTATCTTGGCTACCAAGAGCTTGGGATGGGCAGG
HC2-80	-----
HC2B	-----
HC2C	-----
HC2D-KIAA1058	CCGGTCTCGGCGAACCTTCCTTCGGGCTATCTTGGCTACCAGGAGCTTGGGATGGGCAGG
HC2E	-----
HC2F	-----

HC2A	CATTATGGTCCGGAAATTAAATGGGTAGATGGAGGCAAGCCACTGCTGAAAATTTCCACT
HC2-80	-----
HC2B	-----
HC2C	-----
HC2D-KIAA1058	CATTATGGTCCGGAAATTAAATGGGTAGATGGAGGCAAGCCACTGCTGAAAATTTCCACT
HC2E	-----
HC2F	-----

HC2A	CATCTGGTTTCTACAGTGTATACTCAGGATCAGCATTTACATAATTTTTTCCAGTACTGT
HC2-80	-----
HC2B	-----
HC2C	-----
HC2D-KIAA1058	CATCTGGTTTCTACAGTGTATACTCAGGATCAGCATTTACATAATTTTTTCCAGTACTGT
HC2E	-----
HC2F	-----

HC2A	CAGAAAACCGAATCTGGAGCCCAAGCCTTAGGAAACGAACTTGTAAGTACCTTAAGAGT
HC2-80	-----
HC2B	-----
HC2C	-----
HC2D-KIAA1058	CAGAAAACCGAATCTGGAGCCCAAGCCTTAGGAAACGAACTTGTAAGTACCTTAAGAGT
HC2E	-----
HC2F	-----

HC2A	CTGCATGCGATGGAAGGCCACGTGATGATCGCCTTCTTGCCCACTATCCTAAACCAGCTG
HC2-80	-----
HC2B	-----GCGATGGAAGGCCACGTGATGATCGCCTTCTTGCCCACTATCCTAAACCAGCTG
HC2C	-----
HC2D-KIAA1058	CTGCATGCGATGGAAGGCCACGTGATGATCGCCTTCTTGCCCACTATCCTAAACCAGCTG
HC2E	-----GCGATGGAAGGCCACGTGATGATCGCCTTCTTGCCCACTATCCTAAACCAGCTG
HC2F	-----

Fig. 3A (cont.)

HC2A	TTCCGAGTCCTCACCAGAGCCACACAGGAAGAAGTCGCGGTTAACGTGACTCGGGTCATT
HC2-80	-----
HC2B	TTCCGAGTCCTCACCAGAGCCACACAGGAAGAAGTCGCGGTTAACGTGACTCGGGTCATT
HC2C	-----
HC2D-KIAA1058	TTCCGAGTCCTCACCAGAGCCACACAGGAAGAAGTCGCGGTTAACGTGACTCGGGTCATT
HC2E	TTCCGAGTCCTCACCAGAGCCACACAGGAAGAAGTCGCGGTTAACGTGACTCGGGTCATT
HC2F	-----
HC2A	ATTCATGTGGTTGCCCAGTGCCATGAGGAAGGATTGGAGAGCCACTTGAGGTCATATGTT
HC2-80	-----
HC2B	ATTCATGTGGTTGCCCAGTGCCATGAGGAAGGATTGGAGAGCCACTTGAGGTCATATGTT
HC2C	-----
HC2D-KIAA1058	ATTCATGTGGTTGCCCAGTGCCATGAGGAAGGATTGGAGAGCCACTTGAGGTCATATGTT
HC2E	ATTCATGTGGTTGCCCAGTGCCATGAGGAAGGATTGGAGAGCCACTTGAGGTCATATGTT
HC2F	-----
HC2A	AAGTACGCGTATAAGGCTGAGCCATATGTTGCCTCTGAATACAAGACAGTGCATGAAGAA
HC2-80	-----
HC2B	AAGTACGCGTATAAGGCTGAGCCATATGTTGCCTCTGAATACAAGACAGTGCATGAAGAA
HC2C	-----
HC2D-KIAA1058	AAGTACGCGTATAAGGCTGAGCCATATGTTGCCTCTGAATACAAGACAGTGCATGAAGAA
HC2E	AAGTACGCGTATAAGGCTGAGCCATATGTTGCCTCTGAATACAAGACAGTGCATGAAGAA
HC2F	-----
HC2A	CTGACCAAATCCATGACCACGATTCTCAAGCCTTCTGCCGATTTCTCACCAGCAACAAA
HC2-80	-----
HC2B	CTGACCAAATCCATGACCACGATTCTCAAGCCTTCTGCCGATTTCTCACCAGCAACAAA
HC2C	-----
HC2D-KIAA1058	CTGACCAAATCCATGACCACGATTCTCAAGCCTTCTGCCGATTTCTCACCAGCAACAAA
HC2E	CTGACCAAATCCATGACCACGATTCTCAAGCCTTCTGCCGATTTCTCACCAGCAACAAA
HC2F	-----
HC2A	CTACTGAGGTACTCATGGTTTTTCTTTGATGTACTGATCAAATCTATGGCTCAGCATTTG
HC2-80	-----
HC2B	CTACTGAGGTACTCATGGTTTTTCTTTGATGTACTGATCAAATCTATGGCTCAGCATTTG
HC2C	-----
HC2D-KIAA1058	CTACTGAAGTACTCATGGTTTTTCTTTGATGTACTGATCAAATCTATGGCTCAGCATTTG
HC2E	CTACTGAGGTACTCATGGTTTTTCTTTGATGTACTGATCAAATCTATGGCTCAGCATTTG
HC2F	-----
HC2A	ATAGAGAACTCCAAAGTTAAGTTGCTGCGAAACCAGAGATTTCTGCATCCTATCATCAT
HC2-80	-----
HC2B	ATAGAGAACTCCAAAGTTAAGTTGCTGCGAAACCAGAGATTTCTGCATCCTATCATCAT
HC2C	-----
HC2D-KIAA1058	ATAGAGAACTCCAAAGTTAAGTTGCTGCGAAACCAGAGATTTCTGCATCCTATCATCAT
HC2E	ATAGAGAACTCCAAAGTTAAGTTGCTGCGAAACCAGAGATTTCTGCATCCTATCATCAT
HC2F	-----

Fig. 3A (cont.)

HC2A	GCAGCGGAAACCGTTGTAAATATGCTGATGCCACACATCACTCAGAAGTTTGGAGATAAT
HC2-80	-----
HC2B	GCAGCGGAAACCGTTGTAAATATGCTGATGCCACACATCACTCAGAAGTTTGGAGATAAT
HC2C	-----
HC2D-KIAA1058	GCAGTGGAAACCGTTGTAAATATGCTGATGCCACACATCACTCAGAAGTTTCGAGATAAT
HC2E	GCAGCGGAAACCGTTGTAAATATGCTGATGCCACACATCACTCAGAAGTTTGGAGATAAT
HC2F	-----
HC2A	CCAGAGGCATCTAAGAACGCGAATCATAGCCTTGCTGTCTTCATCAAGAGATGTTTCACC
HC2-80	-----
HC2B	CCAGAGGCATCTAAGAACGCGAATCATAGCCTTGCTGTCTTCATCAAGAGATGTTTCACC
HC2C	-----
HC2D-KIAA1058	CCAGAGGCATCTAAGAACGCGAATCATAGCCTTGCTGTCTTCATCAAGAGATGTTTCACC
HC2E	CCAGAGGCATCTAAGAACGCGAATCATAGCCTTGCTGTCTTCATCAAGAGATGTTTCACC
HC2F	-----
HC2A	TTCATGGACAGGGGCTTTGTCTTCAAGCAGATCAACAACACTACATTAGCTGTTTTGCTCCT
HC2-80	-----
HC2B	TTCATGGACAGGGGCTTTGTCTTCAAGCAGATCAACAACACTACATTAGCTGTTTTGCTCCT
HC2C	-----
HC2D-KIAA1058	TTCATGGACAGGGGCTTTGTCTTCAAGCAGATCAACAACACTACATTAGCTGTTTTGCTCCT
HC2E	TTCATGGACAGGGGCTTTGTCTTCAAGCAGATCAACAACACTACATTAGCTGTTTTGCTCCT
HC2F	-----
HC2A	GGAGACCCAAAGACCCTCTTTGAATACAAGTTTGAATTTCTCCGTGTAGTGTGCAACCAT
HC2-80	-----
HC2B	GGAGACCCAAAGACCCTCTTTGAATACAAGTTTGAATTTCTCCGTGTAGTGTGCAACCAT
HC2C	-----
HC2D-KIAA1058	GGAGACCCAAAGACCCTCTTTGAATACAAGTTTGAATTTCTCCGTGTAGTGTGCAACCAT
HC2E	GGAGACCCAAAGACCCTCTTTGAATACAAGTTTGAATTTCTCCGTGTAGTGTGCAACCAT
HC2F	-----
HC2A	GAACATTATATTCCGTTGAACTTACCAATGCCATTTGGAAAAGGCAGGATTCAAAGATAC
HC2-80	-----
HC2B	GAACATTATATTCCGTTGAACTTACCAATGCCATTTGGAAAAGGCAGGATTCAAAGATAC
HC2C	-----
HC2D-KIAA1058	GAACATTATATTCCGTTGAACTTACCAATGCCATTTGGAAAAGGCAGGATTCAAAGATAC
HC2E	GAACATTATATTCCGTTGAACTTACCAATGCCATTTGGAAAAGGCAGGATTCAAAGATAC
HC2F	-----
HC2A	CAAGACCTCCAGCTTGACTACTCATTAAACAGATGAGTTCTGCAGAAACCACTTCTTGGTG
HC2-80	-----TCCAGCTTGACTACTCATTAAACAGATGAGTTCTGCAGAAACCACTTCTTGGTG
HC2B	CAAGACCTCCAGCTTGACTACTCATTAAACAGATGAGTTCTGCAGAAACCACTTCTTGGTG
HC2C	-----
HC2D-KIAA1058	CAAGACCTCCAGCTTGACTACTCATTAAACAGATGAGTTCTGCAGAAACCACTTCTTGGTG
HC2E	CAAGACCTCCAGCTTGACTACTCATTAAACAGATGAGTTCTGCAGAAACCACTTCTTGGTG
HC2F	-----

FIG. 3A (cont.)

HC2A	GGACTGTTACTGAGGGAGGTGGGGACAGCCCTCCAGGAGTTCCGGGAGGTCCGTCTGATC
HC2-80	GGACTGTTACTGAGGGAGGTGGGGACAGCCCTCCAGGAGTTCCGGGAGGTCCGTCTGATC
HC2B	GGACTGTTACTGAGGGAGGTGGGGACAGCCCTCCAGGAGTTCCGGGAGGTCCGTCTGATC
HC2C	-----
HC2D-KIAA1058	GGACTGTTACTGAGGGAGGTGGGGACAGCCCTCCAGGAGTTCCGGGAGGTCCGTCTGATC
HC2E	GGACTGTTACTGAGGGAGGTGGGGACAGCCCTCCAGGAGTTCCGGGAGGTCCGTCTGATC
HC2F	-----
HC2A	GCCATCAGTGTGCTCAAGAACCTGCTGATAAAGCATTCTTTTGATGACAGATATGCTTCA
HC2-80	GCCATCAGTGTGCTCAAGAACCTGCTGATAAAGCATTCTTTTGATGACAGATATGCTTCA
HC2B	GCCATCAGTGTGCTCAAGAACCTGCTGATAAAGCATTCTTTTGATGACAGATATGCTTCA
HC2C	-----
HC2D-KIAA1058	GCCATCAGTGTGCTCAAGAACCTGCTGATAAAGCATTCTTTTGATGACAGATATGCTTCA
HC2E	GCCATCAGTGTGCTCAAGAACCTGCTGATAAAGCATTCTTTTGATGACAGATATGCTTCA
HC2F	-----
HC2A	AGGAGCCATCAGGCAAGGATAGCCACCCTCTACCTGCCTCTGTTTGGTCTGCTGATTGAA
HC2-80	AGGAGCCATCAGGCAAGGATAGCCACCCTCTACCTGCCTCTGTTTGGTCTGCTGATTGAA
HC2B	AGGAGCCATCAGGCAAGGATAGCCACCCTCTACCTGCCTCTGTTTGGTCTGCTGATTGAA
HC2C	-----
HC2D-KIAA1058	AGGAGCCATCAGGCAAGGATAGCCACCCTCTACCTGCCTCTGTTTGGTCTGCTGATTGAA
HC2E	AGGAGCCATCAGGCAAGGATAGCCACCCTCTACCTGCCTCTGTTTGGTCTGCTGATTGAA
HC2F	-----
HC2A	AACGTCCAGCGGATCAATGTGAGGGATGTGTCACCCTTCCCTGTGAACGCGGGCATGACC
HC2-80	AACGTCCAGCGGATCAATGTGAGGGATGTGTCACCCTTCCCTGTGAACGCGGGCATGACC
HC2B	AACGTCCAGCGGATCAATGTGAGGGATGTGTCACCCTTCCCTGTGAACGCGGGCATGACC
HC2C	-----
HC2D-KIAA1058	AACGTCCAGCGGATCAATGTGAGGGATGTGTCACCCTTCCCTGTGAACGCGGGCATGACT
HC2E	AACGTCCAGCGGATCAATGTGAGGGATGTGTCACCCTTCCCTGTGAACGCGGGCATGACC
HC2F	-----
HC2A	GTGAAGGATGAATCCCTGGCTCTACCAGCTGTGAATCCGCTGGTGACGCCGCAGAAGGGA
HC2-80	GTGAAGGATGAATCCCTGGCTCTACCAGCTGTGAATCCGCTGGTGACGCCGCAGAAGGGA
HC2B	GTGAAGGATGAATCCCTGGCTCTACCAGCTGTGAATCCGCTGGTGACGCCGCAGAAGGGA
HC2C	-----
HC2D-KIAA1058	GTGAAGGATGAATCCCTGGCTCTACCAGCTGTGAATCCGCTGGTGACGCCGCAGAAGGGA
HC2E	GTGAAGGATGAATCCCTGGCTCTACCAGCTGTGAATCCGCTGGTGACGCCGCAGAAGGGA
HC2F	-----
HC2A	AGCACCCCTGGACAACAGCCTGCACAAGGACCTGCTGGGCGCCATCTCCGGCATTGCTTCT
HC2-80	AGCACCCCTGGACAACAGCCTGCACAAGGACCTGCTGGGCGCCATCTCCGGCATTGCTTCT
HC2B	AGCACCCCTGGACAACAGCCTGCACAAGGACCTGCTGGGCGCCATCTCCGGCATTGCTTCT
HC2C	-----
HC2D-KIAA1058	AGCACCCCTGGACAACAGCCTGCACAAGGACCTGCTGGGCGCCATCTCCGGCATTGCTTCT
HC2E	AGCACCCCTGGACAACAGCCTGCACAAGGACCTGCTGGGCGCCATCTCCGGCATTGCTTCT
HC2F	-----

Fig. 3A (cont.)



HC2A	CCATATACAACCTCAACTCCAAACATCAACAGTGTGAGAAATGCTGATTTCGAGAGGATCT
HC2-80	CCATATACAACCTCAACTCCAAACATCAACAGTGTGAGAAATGCTGATTTCGAGAGGATCT
HC2B	CCATATACAACCTCAACTCCAAACATCAACAGTGTGAGAAATGCTGATTTCGAGAGGATCT
HC2C	-----
HC2D-KIAA1058	CCATATACAACCTCAACTCCAAACATCAACAGTGTGAGAAATGCTGATTTCGAGAGGATCT
HC2E	CCATATACAACCTCAACTCCAAACATCAACAGTGTGAGAAATGCTGATTTCGAGAGGATCT
HC2F	-----GCTGATTTCGAGAGGATCT
HC2A	CTCATAAGCACAGATTTCGGGTAACAGCCTTCCAGAAAGGAATAGTGAGAAGAGCAATTCC
HC2-80	CTCATAAGCACAGATTTCGGGTAACAGCCTTCCAGAAAGGAATAGTGAGAAGAGCAATTCC
HC2B	CTCATAAGCACAGATTTCGGGTAACAGCCTTCCAGAAAGGAATAGTGAGAAGAGCAATTCC
HC2C	-----
HC2D-KIAA1058	CTCATAAGCACAGATTTCGGGTAACAGCCTTCCAGAAAGGAATAGTGAGAAGAGCAATTCC
HC2E	CTCATAAGCACAGATTTCGGGTAACAGCCTTCCAGAAAGGAATAGTGAGAAGAGCAATTCC
HC2F	CTCATAAGCACAGATTTCGGGTAACAGCCTTCCAGAAAGGAATAGTGAGAAGAGCAATTCC
HC2A	CTGGATAAGCACCAACAAAGTAGCACATTGGGAAATTCCGTGGTTCGCTGTGATAAACTT
HC2-80	CTGGATAAGCACCAACAAAGTAGCACATTGGGAAATTCCGTGGTTCGCTGTGATAAACTT
HC2B	CTGGATAAGCACCAACAAAGTAGCACATTGGGAAATTCCGTGGTTCGCTGTGATAAACTT
HC2C	-----
HC2D-KIAA1058	CTGGATAAGCACCAACAAAGTAGCACATTGGGAAATTCCGTGGTTCGCTGTGATAAACTT
HC2E	CTGGATAAGCACCAACAAAGTAGCACATTGGGAAATTCCGTGGTTCGCTGTGATAAACTT
HC2F	CTGGATAAGCACCAACAAAGTAGCACATTGGGAAATTCCGTGGTTCGCTGTGATAAACTT
HC2A	GACCAGTCTGAGATTAAGAGCCTACTGATGTGTTTCCTCTACATCTTAAAGAGCATGTCT
HC2-80	GACCAGTCTGAGATTAAGAGCCTACTGATGTGTTTCCTCTACATCTTAAAGAGCATGTCT
HC2B	GACCAGTCTGAGATTAAGAGCCTACTGATGTGTTTCCTCTACATCTTAAAGAGCATGTCT
HC2C	-----
HC2D-KIAA1058	GACCAGTCTGAGATTAAGAGCCTACTGATGTGTTTCCTCTACATCTTAAAGAGCATGTCT
HC2E	GACCAGTCTGAGATTAAGAGCCTACTGATGTGTTTCCTCTACATCTTAAAGAGCATGTCT
HC2F	GACCAGTCTGAGATTAAGAGCCTACTGATGTGTTTCCTCTACATCTTAAAGAGCATGTCT
HC2A	GATGATGCTTTGTTTACATATTGGAACAAGGCTTCAACATCTGAACTTATGGATTTTTTTT
HC2-80	GATGATGCTTTGTTTACATATTGGAACAAGGCTTCAACATCTGAACTTATGGATTTTTTTT
HC2B	GATGATGCTTTGTTTACATATTGGAACAAGGCTTCAACATCTGAACTTATGGATTTTTTTT
HC2C	-----
HC2D-KIAA1058	GATGATGCTTTGTTTACATATTGGAACAAGGCTTCAACATCTGAACTTATGGATTTTTTTT
HC2E	GATGATGCTTTGTTTACATATTGGAACAAGGCTTCAACATCTGAACTTATGGATTTTTTTT
HC2F	GATGATGCTTTGTTTACATATTGGAACAAGGCTTCAACATCTGAACTTATGGATTTTTTTT
HC2A	ACAATATCTGAAGTCTGCCTGCACCAGTTCCAGTACATGGGGAAGCGATACATAGCCAGG
HC2-80	ACAATATCTGAAGTCTGCCTGCACCAGTTCCAGTACATGGGGAAGCGATACATAGCCAGG
HC2B	ACAATATCTGAAGTCTGCCTGCACCAGTTCCAGTACATGGGGAAGCGATACATAGCCAGG
HC2C	-----
HC2D-KIAA1058	ACAATATCTGAAGTCTGCCTGCACCAGTTCCAGTACATGGGGAAGCGATACATAGCCAG-
HC2E	ACAATATCTGAAGTCTGCCTGCACCAGTTCCAGTACATGGGGAAGCGATACATAGCCAGG
HC2F	ACAATATCTGAAGTCTGCCTGCACCAGTTCCAGTACATGGGGAAGCGATACATAGCCAG-

Fig. 3A (cont.)

HC2A	AACCAGGAGGGGTTGGGACCCATAGTTCATGATCGAAAGTCTCAGACATTGCCTGTTTCC
HC2-80	AACCAGGAGGGGTTGGGACCCATAGTTCATGATCGAAAGTCTCAGACATTGCCTGTTTCC
HC2B	AACCAGGAGGGGTTGGGACCCATAGTTCATGATCGAAAGTCTCAGACATTGCCTGTTTCC
HC2C	-----
HC2D-KIAA1058	-----AA-----
HC2E	AACCAGGAGGGGTTGGGACCCATAGTTCATGATCGAAAGTCTCAGACATTGCCTGTTTCC
HC2F	-----TGTGA-----GAAAG-----ATATCAAGTGT-----
HC2A	CGTAACAGAACAGGAATGATGCATGCCAGATTGCAGCAGCTGGGCAGCCTGGATAACTCT
HC2-80	CGTAACAGAACAGGAATGATGCATGCCAGATTGCAGCAGCTGGGCAGCCTGGATAACTCT
HC2B	CGTAACAGAACAGGAATGATGCATGCCAGATTGCAGCAGCTGGGCAGCCTGGATAACTCT
HC2C	-----
HC2D-KIAA1058	-----CAGGAATGATGCATGCCAGATTGCAGCAGCTGGGCAGCCTGGATAACTCT
HC2E	CGTAACAGAACAGGAATGATGCATGCCAGATTGCAGCAGCTGGGCAGCCTGGATAACTCT
HC2F	-----GCTTGGAA-----
HC2A	CTCACTTTTAACACAGCTATGGCCACTCGGACGCAGATGTTCTGCACCAGTCATTACTT
HC2-80	CTCACTTTTAACACAGCTATGGCCACTCGGACGCAGATGTTCTGCACCAGTCATTACTT
HC2B	CTCACTTTTAACACAGCTATGGCCACTCGGACGCAGATGTTCTGCACCAGTCATTACTT
HC2C	-----
HC2D-KIAA1058	CTCACTTTTAACACAGCTATGGCCACTCGGACGCAGATGTTCTGCACCAGTCATTACTT
HC2E	CTCACTTTTAACACAGCTATGGCCACTCGGACGCAGATGTTCTGCACCAGTCATTACTT
HC2F	-TTTCTGTAGACAATGGCTATGGCCACTCGGACGCAGATGTTCTGCACCAGTCATTACTT
HC2A	GAAGCCAACATTGCTACTGAGGTTTGCCTGACAGCTCTGGACACGCTTTCTCTATTTACA
HC2-80	GAAGCCAACATTGCTACTGAGGTTTGCCTGACAGCTCTGGACACGCTTTCTCTATTTACA
HC2B	GAAGCCAACATTGCTACTGAGGTTTGCCTGACAGCTCTGGACACGCTTTCTCTATTTACA
HC2C	-----
HC2D-KIAA1058	GAAGCCAACATTGCTACTGAGGTTTGCCTGACAGCTCTGGACACGCTTTCTCTATTTACA
HC2E	GAAGCCAACATTGCTACTGAGGTTTGCCTGACAGCTCTGGACACGCTTTCTCTATTTACA
HC2F	GAAGCCAACATTGCTACTGAGGTTTGCCTGACAGCTCTGGACACGCTTTCTCTATTTACA
HC2A	TTGGCGTTTAAGAACCAGCTCCTGGCCGACCATGGACATAATCCTCTCATGAAAAAAGTT
HC2-80	TTGGCGTTTAAGAACCAGCTCCTGGCCGACCATGGACATAATCCTCTCATGAAAAAAGTT
HC2B	TTGGCGTTTAAG-----CTCCTGGCCGACCATGGACATAATCCTCTCATGAAAAAAGTT
HC2C	-----
HC2D-KIAA1058	TTGGCGTTTAAGAACCAGCTCCTGGCCGACCATGGACATAATCCTCTCATGAAAAAAGTT
HC2E	TTGGCGTTTAAGAACCAGCTCCTGGCCGACCATGGACATAATCCTCTCATGAAAAAAGTT
HC2F	TTGGCGTTTAAGAACCAGCTCCTGGCCGACCATGGACATAATCCTCTCATGAAAAAAGTT
HC2A	TTTGATGTCTACCTGTGTTTTCTTCAAAAACATCAGTCTGAAACGGCTTTAAAAAATGTC
HC2-80	TTTGATGTCTACCTGTGTTTTCTTCAAAAACATCAGTCTGAAACGGCTTTAAAAAATGTC
HC2B	TTTGATGTCTACCTGTGTTTTCTTCAAAAACATCAGTCTGAAACGGCTTTAAAAAATGTC
HC2C	-----
HC2D-KIAA1058	TTTGATGTCTACCTGTGTTTTCTTCAAAAACATCAGTCTGAAACGGCTTTAAAAAATGTC
HC2E	TTTGATGTCTACCTGTGTTTTCTTCAAAAACATCAGTCTGAAACGGCTTTAAAAAATGTC
HC2F	A-----

Fig. 3A (cont.)

HC2A  
HC2-80  
HC2B  
HC2C  
HC2D-KIAA1058  
HC2E  
HC2F

TTCACTGCCTTAAGGTCCTTAATTTATAAGTTTCCCTCAACATTCTATGAAGGGAGAGCG  
TTCACTGCCTTAAGGTCCTTAATTTATAAGTTTCCCTCAACATTCTATGAAGGGAGAGCG  
TTCACTGCCTTAAGGTCCTTAATTTATAAGTTTCCCTCAACATTCTATGAAGGGAGAGCG  
-----  
TTCACTGCCTTAAGGTCCTTAATTTATAAGTTTCCCTCAACATTCTATGAAGGGAGAGCG  
TTCACTGCCTTAAGGTCCTTAATTTATAAGTTTCCCTCAACATTCTATGAAGGGAGAGCG  
-----

HC2A  
HC2-80  
HC2B  
HC2C  
HC2D-KIAA1058  
HC2E  
HC2F

GACATGTGTGCGGCTCTGTGTTACGAGATTCTCAAGTGCTGTAACCTCCAAGCTGAGCTCC  
GACATGTGTGCGGCTCTGTGTTACGAGATTCTCAAGTGCTGTAACCTCCAAGCTGAGCTCC  
GACATGTGTGCGGCTCTGTGTTACGAGATTCTCAAGTGCTGTAACCTCCAAGCTGAGCTCC  
-----  
GACATGTGTGCGGCTCTGTGTTACGAGATTCTCAAGTGCTGTAACCTCCAAGCTGAGCTCC  
GACATGTGTGCGGCTCTGTGTTACGAGATTCTCAAGTGCTGTAACCTCCAAGCTGAGCTCC  
-----

HC2A  
HC2-80  
HC2B  
HC2C  
HC2D-KIAA1058  
HC2E  
HC2F

ATCAGGACGGAGGCCTCCCAGCTGCTCTACTTCCTGATGAGGAACAACCTTTGATTACACT  
ATCAGGACGGAGGCCTCCCAGCTGCTCTACTTCCTGATGAGGAACAACCTTTGATTACACT  
ATCAGGACGGAGGCCTCCCAGCTGCTCTACTTCCTGATGAGGAACAACCTTTGATTACACT  
-----  
ATCAGGACGGAGGCCTCCCAGCTGCTCTACTTCCTGATGAGGAACAACCTTTGATTACACT  
ATCAGGACGGAGGCCTCCCAGCTGCTCTACTTCCTGATGAGGAACAACCTTTGATTACACT  
-----

HC2A  
HC2-80  
HC2B  
HC2C  
HC2D-KIAA1058  
HC2E  
HC2F

GGAAAGAAGTCCTTTGTCCGGACACATTTGCAAGTCATCATATCTGTCAGCCAGCTGATA  
GGAAAGAAGTCCTTTGTCCGGACACATTTGCAAGTCATCATATCTGTCAGCCAGCTGATA  
GGAAAGAAGTCCTTTGTCCGGACACATTTGCAAGTCATCATATCTGTCAGCCAGCTGATA  
-----  
GGAAAGAAGTCCTTTGTCCGGACACATTTGCAAGTCATCATATCTGTCAGCCAGCTGATA  
GGAAAGAAGTCCTTTGTCCGGACACATTTGCAAGTCATCATATCTGTCAGCCAGCTGATA  
-----

HC2A  
HC2-80  
HC2B  
HC2C  
HC2D-KIAA1058  
HC2E  
HC2F

GCAGACGTTGTTGGCATTGGGGAAACCAGATTCCAGCAGTCCCTGTCCATCATCAACAAC  
GCAGACGTTGTTGGCATTGGGGAAACCAGATTCCAGCAGTCCCTGTCCATCATCAACAAC  
GCAGACGTTGTTGGCATTGGGGAAACCAGATTCCAGCAGTCCCTGTCCATCATCAACAAC  
-----  
GCAGACGTTGTTGGCATTGGGGAAACCAGATTCCAGCAGTCCCTGTCCATCATCAACAAC  
GCAGACGTTGTTGGCATTGGGGAAACCAGATTCCAGCAGTCCCTGTCCATCATCAACAAC  
-----

HC2A  
HC2-80  
HC2B  
HC2C  
HC2D-KIAA1058  
HC2E  
HC2F

TGTGCCAACAGTGACCGGCTTATTAAGCACACCAGCTTCTCCTCTGATGTGAAGGACTTA  
TGTGCCAACAGTGACCGGCTTATTAAGCACACCAGCTTCTCCTCTGATGTGAAGGACTTA  
TGTGCCAACAGTGACCGGCTTATTAAGCACACCAGCTTCTCCTCTGATGTGAAGGACTTA  
-----  
TGTGCCAACAGTGACCGGCTTATTAAGCACACCAGCTTCTCCTCTGATGTGAAGGACTTA  
TGTGCCAACAGTGACCGGCTTATTAAGCACACCAGCTTCTCCTCTGATGTGAAGGACTTA  
-----

FIG. 3A (cont.)

HC2A	ACCAAAAGGATACGCACGGTGCTAATGGCCACCGCCCAGATGAAGGAGCATGAGAACGAC
HC2-80	ACCAAAAGGATACGCACGGTGCTAATGGCCACCGCCCAGATGAAGGAGCATGAGAACGAC
HC2B	ACCAAAAGGATACGCACGGTGCTAATGGCCACCGCCCAGATGAAGGAGCATGAGAACGAC
HC2C	-----
HC2D-KIAA1058	ACCAAAAGGATACGCACGGTGCTAATGGCCACCGCCCAGATGAAGGAGCATGAGAACGAC
HC2E	ACCAAAAGGATACGCACGGTGCTAATGGCCACCGCCCAGATGAAGGAGCATGAGAACGAC
HC2F	-----
HC2A	CCAGAGATGCTGGTGGACCTCCAGTACAGCCTGGCCAAATCCTATGCCAGCACGCCCCGAG
HC2-80	CCAGAGATGCTGGTGGACCTCCAGTACAGCCTGGCCAAATCCTATGCCAGCACGCCCCGAG
HC2B	CCAGAGATGCTGGTGGACCTCCAGTACAGCCTGGCCAAATCCTATGCCAGCACGCCCCGAG
HC2C	-----
HC2D-KIAA1058	CCAGAGATGCTGGTGGACCTCCAGTACAGCCTGGCCAAATCCTATGCCAGCACGCCCCGAG
HC2E	CCAGAGATGCTGGTGGACCTCCAGTACAGCCTGGCCAAATCCTATGCCAGCACGCCCCGAG
HC2F	-----
HC2A	CTCAGGAAGACGTGGCTCGACAGCATGGCCAGGATCCATGTCAAAAATGGCGATCTCTCA
HC2-80	CTCAGGAAGACGTGGCTCGACAGCATGGCCAGGATCCATGTCAAAAATGGCGATCTCTCA
HC2B	CTCAGGAAGACGTGGCTCGACAGCATGGCCAGGATCCATGTCAAAAATGGCGATCTCTCA
HC2C	-----
HC2D-KIAA1058	CTCAGGAAGACGTGGCTCGACAGCATGGCCAGGATCCATGTCAAAAATGGCGATCTCTCA
HC2E	CTCAGGAAGACGTGGCTCGACAGCATGGCCAGGATCCATGTCAAAAATGGCGATCTCTCA
HC2F	-----
HC2A	GAGGCAGCAATGTGCTATGTCCACGTAACAGCCCTAGTGGCAGAATATCTCACACGGAAA
HC2-80	GAGGCAGCAATGTGCTATGTCCACGTAACAGCCCTAGTGGCAGAATATCTCACACGGAAA
HC2B	GAGGCAGCAATGTGCTATGTCCACGTAACAGCCCTAGTGGCAGAATATCTCACACGGAAA
HC2C	-----
HC2D-KIAA1058	GAGGCAGCAATGTGCTATGTCCACGTAACAGCCCTAGTGGCAGAATATCTCACACGGAAA
HC2E	GAGGCAGCAATGTGCTATGTCCACGTAACAGCCCTAGTGGCAGAATATCTCACACGGAAA
HC2F	-----
HC2A	G-----
HC2-80	G-----
HC2B	G-----
HC2C	-----
HC2D-KIAA1058	GAAGCAGTCCAGTGGGAGCCGCCCTTCTCCCCACAGCCATAGCGCCTGCCTGAGGAGG
HC2E	G-----
HC2F	-----
HC2A	-----GCGTGTTTAGACAAGGATGCACCGCCTTCAGGGTCATTACCCCAAACATC
HC2-80	-----GCGTGTTTAGACAAGGATGCACCGCCTTCAGGGTCATTACCCCAAACATC
HC2B	-----GCGTGTTTAGACAAGGATGCACCGCCTTCAGGGTCATTACCCCAAACATC
HC2C	-----GTTTAGACAAGGATGCACCGCCTTCAGGGTCATTACCCCAAACATC
HC2D-KIAA1058	AGCCGGGGAGGCGTGTTTAGACAAGGATGCACCGCCTTCAGGGTCATTACCCCAAACATC
HC2E	-----GCGTGTTTAGACAAGGATGCACCGCCTTCAGGGTCATTACCCCAAACATC
HC2F	-----

FIG. 3A (cont.)

HC2A  
HC2-80  
HC2B  
HC2C  
HC2D-KIAA1058  
HC2E  
HC2F

GACGAGGAGGCCTCCATGATGGAAGACGTGGGGATGCAGGATGTCCATTTCAACGAGGAT  
GACGAGGAGGCCTCCATGATGGAAGACGTGGGGATGCAGGATGTCCATTTCAACGAGGAT  
GACGAGGAGGCCTCCATGATGGAAGACGTGGGGATGCAGGATGTCCATTTCAACGAGGAT  
GACGAGGAGGCCTCCATGATGGAAGACGTGGGGATGCAGGATGTCCATTTCAACGAGGAT  
GACGAGGAGGCCTCCATGATGGAAGACGTGGGGATGCAGGATGTCCATTTCAACGAGGAT  
GACGAGGAGGCCTCCATGATGGAAGACGTGGGGA-----  
-----

HC2A  
HC2-80  
HC2B  
HC2C  
HC2D-KIAA1058  
HC2E  
HC2F

GTGCTGATGGAGCTCCTTGAGCAGTGCGCAGATGGACTCTGGAAAGCCGAGCGCTACGAG  
GTGCTGATGGAGCTCCTTGAGCAGTGCGCAGATGGACTCTGGAAAGCCGAGCGCTACGAG  
GTGCTGATGGAGCTCCTTGAGCAGTGCGCAGATGGACTCTGGAAAGCCGAGCGCTACGAG  
GTGCTGATGGAGCTCCTTGAGCAGTGCGCAGATGGACTCTGGAAAGCCGAGCGCTACGAG  
GTGCTGATGGAGCTCCTTGAGCAGTGCGCAGATGGACTCTGGAAAGCCGAGCGCTACGAG  
-----AAGCCGAGCGCTACGAG  
-----

HC2A  
HC2-80  
HC2B  
HC2C  
HC2D-KIAA1058  
HC2E  
HC2F

CTCATCGCCGACATCTACAACTTATCATCCCCATTTATGAGAAGCGGAGGGATTT-----  
CTCATCGCCGACATCTACAACTTATCATCCCCATTTATGAGAAGCGGAGGGATTT-----  
CTCATCGCCGACATCTACAACTTATCATCCCCATTTATGAGAAGCGGAGGGATTTTGAG  
CTCATCGCCGACATCTACAACTTATCATCCCCATTTATGAGAAGCGGAGGGATTTTGAG  
CTCATTGCCGACATCTACAACTTATCATCCCCATTTATGAGAAGCGGAGGGATTTTGAG  
CTCATCGCCGACATCTACAACTTATCATCCCCATTTATGAGAAGCGGAGGGATTTTGAG  
-----

HC2A  
HC2-80  
HC2B  
HC2C  
HC2D-KIAA1058  
HC2E  
HC2F

-----  
-----  
AGGCTGGCCCATCTGTATGACACGCTGCACCGGGCCTACAGCAAAGTGACCGAGGTCATG  
AGGCTGGCCCATCTGTATGACACGCTGCACCGGGCCTACAGCAAAGTGACCGAGGTCATG  
AGGCTGGCCCATCTGTATGACACGCTGCACCGGGCCTACAGCAAAGTGACCGAGGTCATG  
AGGCTGGCCCATCTGTATGACACGCTGCACCGGGCCTACAGCAAAGTGACCGAGGTCATG  
-----

HC2A  
HC2-80  
HC2B  
HC2C  
HC2D-KIAA1058  
HC2E  
HC2F

-----  
-----  
CACTCGGGCCGCAGGCTTCTGGGGACCTACTTCCGGGTAGCCTTCTTCGGGCAGG-----  
CACTCGGGCCGCAGGCTTCTGGGGACCTACTTCCGGGTAGCCTTCTTCGGGCAGG-----  
CACTCGGGCCGCAGGCTTCTGGGGACCTACTTCCGGGTAGCCTTCTTCGGGCAGGCAAGC  
CACTCGGGCCGCAGGCTTCTGGGGACCTACTTCCGGGTAGCCTTCTTCGGGCAGG-----  
-----

HC2A  
HC2-80  
HC2B  
HC2C  
HC2D-KIAA1058  
HC2E  
HC2F

-----CTTTGAAGATGAAGATGGA  
-----CTTTGAAGATGAAGATGGA  
-----GATTCTTTGAAGATGAAGATGGA  
-----GATTCTTTGAAGATGAAGATGGA  
CAATACCAGTTTACAGACAGTGAAACAGATGTGGAGGGATTCTTTGAAGATGAAGATGGA  
-----GATTCTTTGAAGATGAAGATGGA  
-----

FIG. 3A (cont.)



0963450

HC2A  
HC2-80  
HC2B  
HC2C  
HC2D-KIAA1058  
HC2E  
HC2F

AAGGAGTATATTTACAAGGAACCCAAACTCACACCGCTGTCGGAAATTTCTCAGAGACTC  
AAGGAGTATATTTACAAGGAACCCAAACTCACACCGCTGTCGGAAATTTCTCAGAGACTC  
AAGGAGTATATTTACAAGGAACCCAAACTCACACCGCTGTCGGAAATTTCTCAGAGACTC  
AAGGAGTATATTTACAAGGAACCCAAACTCACACCGCTGTCGGAAATTTCTCAGAGACTC  
AAGGAGTATATTTACAAGGAACCCAAACTCACACCGCTGTCGGAAATTTCTCAGAGACTC  
AAGGAGTATATTTACAAGGAACCCAAACTCACACCGCTGTCGGAAATTTCTCAGAGACTC  
-----

HC2A  
HC2-80  
HC2B  
HC2C  
HC2D-KIAA1058  
HC2E  
HC2F

CTTAAACTGTACTCGGATAAATTTGGTTCTGAAAATGTCAAAATGATACAGGATTCTGGC  
CTTAAACTGTACTCGGATAAATTTGGTTCTGAAAATGTCAAAATGATACAGGATTCTGGC  
CTTAAACTGTACTCGGATAAATTTGGTTCTGAAAATGTCAAAATGATACAGGATTCTGGC  
CTTAAACTGTACTCGGATAAATTTGGTTCTGAAAATGTCAAAATGATACAGGATTCTGGC  
CTTAAACTGTACTCGGATAAATTTGGTTCTGAAAATGTCAAAATGATACAGGATTCTGGC  
CTTAAACTGTACTCGGATAAATTTGGTTCTGAAAATGTCAAAATGATACAGGATTCTGGC  
-----

HC2A  
HC2-80  
HC2B  
HC2C  
HC2D-KIAA1058  
HC2E  
HC2F

AAGGTCAACCCTAAGGATCTGGATTCTAAGTATGCATACATCCAGGTGACTCACGTCATC  
AAGGTCAACCCTAAGGATCTGGATTCTAAGTATGCATACATCCAGGTGACTCACGTCATC  
AAGGTCAACCCTAAGGATCTGGATTCTAAGTATGCATACATCCAGGTGACTCACGTCATC  
AAGGTCAACCCTAAGGATCTGGATTCTAAGTATGCATACATCCAGGTGACTCACGTCATC  
AAGGTCAACCCTAAGGATCTGGATTCTAAGTATGCCTACATCCAGGTGACTCACGTCATC  
AAGGTCAACCCTAAGGATCTGGATTCTAAGTATGCATACATCCAGGTGACTCACGTCATC  
-----

HC2A  
HC2-80  
HC2B  
HC2C  
HC2D-KIAA1058  
HC2E  
HC2F

CCCTTCTTTGACGAAAAAGAGTTGCAAGAAAGGAAAACAGAGTTTGAGAGATCCCACAAC  
CCCTTCTTTGACGAAAAAGAGTTGCAAGAAAGGAAAACAGAGTTTGAGAGATCCCACAAC  
CCCTTCTTTGACGAAAAAGAGTTGCAAGAAAGGAAAACAGAGTTTGAGAGATCCCACAAC  
CCCTTCTTTGACGAAAAAGAGTTGCAAGAAAGGAAAACAGAGTTTGAGAGATCCCACAAC  
CCCTTCTTTGACGAAAAAGAGTTGCAAGAAAGGAAAACAGAGTTTGAGAGATCCCACAAC  
CCCTTCTTTGACGAAAAAGAGTTGCAAGAAAGGAAAACAGAGTTTGAGAGATCCCACAAC  
-----

HC2A  
HC2-80  
HC2B  
HC2C  
HC2D-KIAA1058  
HC2E  
HC2F

ATCCGCCGCTTCATGTTTGAGATGCCATTTACGCAGACCGGGAAGAGGCAGGGCGGGGTG  
ATCCGCCGCTTCATGTTTGAGATGCCATTTACGCAGACCGGGAAGAGGCAGGGCGGGGTG  
ATCCGCCGCTTCATGTTTGAGATGCCATTTACGCAGACCGGGAAGAGGCAGGGCGGGGTG  
ATCCGCCGCTTCATGTTTGAGATGCCATTTACGCAGACCGGGAAGAGGCAGGGCGGGGTG  
ATCCGCCGCTTCATGTTTGAGATGCCATTTACGCAGACCGGGAAGAGGCAGGGCGGGGTG  
ATCCGCCGCTTCATGTTTGAGATGCCATTTACGCAGACCGGGAAGAGGCAGGGCGGGGTG  
-----

HC2A  
HC2-80  
HC2B  
HC2C  
HC2D-KIAA1058  
HC2E  
HC2F

GAAGAGCAGTGCAAACGGCGCACCATCCTGACAGCCATACTGCTTCCCTTATGTGAAG  
GAAGAGCAGTGCAAACGGCGCACCATCCTGACAGCCATACTGCTTCCCTTATGTGAAG  
GAAGAGCAGTGCAAACGGCGCACCATCCTGACAGCCATACTGCTTCCCTTATGTGAAG  
GAAGAGCAGTGCAAACGGCGCACCATCCTGACAGCCATACTGCTTCCCTTATGTGAAG  
GAAGAGCAGTGCAAACGGCGCACCATCCTGACAGCCATACTGCTTCCCTTATGTGAAG  
GAAGAGCAGTGCAAACGGCGCACCATCCTGACAGCCATACTGCTTCCCTTATGTGAAG  
-----

Fig. 3A (cont.)

HC2A	AAGCGCATCCCTGTCATGTACCAGCACCACACTGACCTGAACCCCATCGAGGTGGCCATT
HC2-80	AAGCGCATCCCTGTCATGTACCAGCACCACACTGACCTGAACCCCATCGAGGTGGCCATT
HC2B	AAGCGCATCCCTGTCATGTACCAGCACCACACTGACCTGAACCCCATCGAGGTGGCCATT
HC2C	AAGCGCATCCCTTTTCATGTACCAGCACCACACTGACCTGAACCCCATCGAGGT--CCATT
HC2D-KIAA1058	AAGCGCATCCCTGTCATGTACCAGCACCACACTGACCTGAACCCCATCGAGGTGGCCATT
HC2E	AAGCGCATCCCTGTCATGTACCAGCACCACACTGACCTGAACCCCATCGAGGTGGCCATT
HC2F	-----

HC2A	GACGAGATGAGTAAGAAGGTGGCGGAGCTCCGGCAGCTGTGCTCCTCGGCCGAGGTGGAC
HC2-80	GACGAGATGAGTAAGAAGGTGGCGGAGCTCCGGCAGCTGTGCTCCTCGGCCGAGGTGGAC
HC2B	GACGAGATGAGTAAGAAGGTGGCGGAGCTCCGGCAGCTGTGCTCCTCGGCCGAGGTGGAC
HC2C	GACGAGATGAGTAAGAAGGTGGCGGAGCTCCGGCAGCTGTGCTCCTCGGCCGAGGTGGAC
HC2D-KIAA1058	GACGAGATGAGTAAGAAGGTGGCGGAGCTCCGGCAGCTGTGCTCCTCGGCCGAGGTGGAC
HC2E	GACGAGATGAGTAAGAAGGTGGCGGAGCTCCGGCAGCTGTGCTCCTCGGCCGAGGTGGAC
HC2F	-----

HC2A	ATGATCAAACCTGCAGCTCAAACCTCCAGGGCAGCGTGAGTGTTTCAGGTCAATGCTGGCCCA
HC2-80	ATGATCAAACCTGCAGCTCAAACCTCCAGGGCAGCGTGAGTGTTTCAGGTCAATGCTGGCCCA
HC2B	ATGATCAAACCTGCAGCTCAAACCTCCAGGGCAGCGTGAGTGTTTCAGGTCAATGCTGGCCCA
HC2C	ATGATCAAACCTGCAGCTCAAACCTCCAGGGCAGCGTGAGTGTTTCAGGTCAATGCTGGCCCA
HC2D-KIAA1058	ATGATCAAACCTGCAGCTCAAACCTCCAGGGCAGCGTGAGTGTTTCAGGTCAATGCTGGCCCA
HC2E	ATGATCAAACCTGCAGCTCAAACCTCCAGGGCAGCGTGAGTGTTTCAGGTCAATGCTGGCCCA
HC2F	-----

HC2A	CTAGCATATGCGCGAGCTTTCTTAGATGATACAAACACAAAGCGATATCCTGACAATAAA
HC2-80	CTAGCATATGCGCGAGCTTTCTTAGATGATACAAACACAAAGCGATATCCTGACAATAAA
HC2B	CTAGCATATGCGCGAGCTTTCTTAGATGATACAAACACAAAGCGATATCCTGACAATAAA
HC2C	CTAGCATATGCGCGAGCTTTCTTAGATGATACAAACACAAAGCGATATCCTGACAATAAA
HC2D-KIAA1058	CTAGCATATGCGCGAGCTTTCTTAGATGATACAAACACAAAGCGATATCCTGACAATAAA
HC2E	CTAGCATATGCGCGAGCTTTCTTAGATGATACAAACACAAAGCGATATCCTGACAATAAA
HC2F	-----

HC2A	GTGAAGCTGCTTAAGGAAGTTTTTCAGGCAATTTGTGGAAGCTTGCGGTCAAGCCTTAGCG
HC2-80	GTGAAGCTGCTTAAGGAAGTTTTTCAGGCAATTTGTGGAAGCTTGCGGTCAAGCCTTAGCG
HC2B	GTGAAGCTGCTTAAGGAAGTTTTTCAGGCAATTTGTGGAAGCTTGCGGTCAAGCCTTAGCG
HC2C	GTGAAGCTGCTTAAGGAAGTTTTTCAGGCAATTTGTGGAAGCTTGCGGTCAAGCCTTAGCG
HC2D-KIAA1058	GTGAAGCTGCTTAAGGAAGTTTTTCAGGCAATTTGTGGAAGCTTGCGGTCAAGCCTTAGCG
HC2E	GTGAAGCTGCTTAAGGAAGTTTTTCAGGCAATTTGTGGAAGCTTGCGGTCAAGCCTTAGCG
HC2F	-----

HC2A	GTAAACGAACGTCTGATTAAAGAAGACCAGCTCGAGTATCAGGAAGAAATGAAAGCCAAC
HC2-80	GTAAACGAACGTCTGATTAAAGAAGACCAGCTCGAGTATCAGGAAGAAATGAAAGCCAAC
HC2B	GTAAACGAACGTCTGATTAAAGAAGACCAGCTCGAGTATCAGGAAGAAATGAAAGCCAAC
HC2C	GTAAACGAACGTCTGATTAAAGAAGACCAGCTCGAGTATCAGGAAGAAATGAAAGCCAAC
HC2D-KIAA1058	GTAAACGAACGTCTGATTAAAGAAGACCAGCTCGAGTATCAGGAAGAAATGAAAGCCAAC
HC2E	GTAAACGAACGTCTGATTAAAGAAGACCAGCTCGAGTATCAGGAAGAAATGAAAGCCAAC
HC2F	-----

Fig. 3A (cont.)

HC2A	TACAGGGAAATGGCGAAGGAGCTTTCTGAAATCATGCATGAGCAG-----ATCTGCC
HC2-80	TACAGGGAAATGGCGAAGGAGCTTTCTGAAATCATGCATGAGCAG-----ATCTGCC
HC2B	TACAGGGAAATGGCGAAGGAGCTTTCTGAAATCATGCATGAGCAG-----ATCTGCC
HC2C	TACAGGGAAATGGCGAAGGAGCTTTCTGAAATCATGCATGAGCAG-----ATCTGCC
HC2D-KIAA1058	TACAGGGAAATGGCGAAGGAGCTTTCTGAAATCATGCATGAGCAGCTGGGATGATCTGCC
HC2E	TACAGGGAAATGGCGAAGGAGCTTTCTGAAATCATGCATGAGCAG-----ATCTGCC
HC2F	-----
HC2A	CCCTGGAGGAGAAGACGAGCGTCTTACCGAATTCCTTCACATCTTCAACGCCATCAGTG
HC2-80	CCCTGGAGGAGAAGACGAGCGTCTTACCGAATTCCTTCACATCTTCAACGCCATCAGTG
HC2B	CCCTGGAGGAGAAGACGAGCGTCTTACCGAATTCCTTCACATCTTCAACGCCATCAGTG
HC2C	CCCTGGAGGAGAAGACGAGCGTCTTACCGAATTCCTTCACATCTTCAACGCCATCAGTG
HC2D-KIAA1058	CCCTGGAGGAGAAGACGAGCGTCTTACCGAATTCCTTCACATCTTCAACGCCATCAGTG
HC2E	CCCTGGAGGAGAAGACGAGCGTCTTACCGAATTCCTTCACATCTTCAACGCCATCAGTG
HC2F	-----
HC2A	GGACTCCAACAAGCACAATGGTTCACGGGATGACCAGCTCGTCTTCGGTCGTGTGATTAC
HC2-80	GGACTCCAACAAGCACAATGGTTCACGGGATGACCAGCTCGTCTTCGGTCGTGTGATTAC
HC2B	GGACTCCAACAAGCACAATGGTTCACGGGATGACCAGCTCGTCTTCGGTCGTGTGA----
HC2C	GGACTCCAACAAGCACAATGGTTCACGGGATGACCAGCTCGTCTTCGGTCGTGTGA----
HC2D-KIAA1058	GGACTCCAACAAGCACAATGGTTCACGGGATGACCAGCTCGTCTTCGGTCGTGTGATTAC
HC2E	GGACTCCAACAAGCACAATGGTTCACGGGATGACCAGCTCGTCTTCGGTCGTGTGA----
HC2F	-----
HC2A	ATCTCATGGCCCGTGTGTGGGGACTTGCTTTGTCAATTTGCAAACCTCAGGATGCTTTCCAA
HC2-80	ATCTCATGGCCCGTGTGTGGGGACTTGCTTTGTCAATTTGCAAACCTCAGGATGCTTTCCAA
HC2B	-----
HC2C	-----
HC2D-KIAA1058	ATCTCATGGCCCGTGTGTGGGGACTTGCTTTGTCAATTTGCAAACCTCAGGATGCTTTCCAA
HC2E	-----
HC2F	-----
HC2A	AGCCAATCACTGGGGAGACCGAGCACAGGGAGGACCAAGGGGAAGGGGAGAGAAAGGAAA
HC2-80	AGCCAATCACTGGGGAGACCGAGCACAGGGAGGACCAAGGGGAAGGGGAGAGAAAGGAAA
HC2B	-----
HC2C	-----
HC2D-KIAA1058	AGCCAATCACTGGGGAGACCGAGCACAGGGAGGACCA-GGGGAAGGGGAGAGAAAGGAAA
HC2E	-----
HC2F	-----
HC2A	TAAAGAACAACGTTATTTCTTAACAGACTTTCTATAGGAGTTGTAAGAAGGTGCACATAT
HC2-80	TAAAGAACAACGTTATTTCTTAACAGACTTTCTATAGGAGTTGTAAGAAGGTGCACATAT
HC2B	-----
HC2C	-----
HC2D-KIAA1058	TAAAGAACAACGTTATTTCTTAACAGACTTTCTATAGGAGTTGTAAGAAGGTGCACATAT
HC2E	-----
HC2F	-----

FIG. 3A (cont.)

HC2A	TTTTTTAAATCTCACTGGCAATATTCAAAGTTTTTCATTGTGTCTTAACAAAGGTGTGGTA
HC2-80	TTTTTTAAATCTCACTGGCAATATTCAAAGTTTTTCATTGTGTCTTAACAAAGGTGTGGTA
HC2B	-----
HC2C	-----
HC2D-KIAA1058	TTTTTTAAATCTCACTGGCAATATTCAAAGTTTTTCATTGTGTCTTAACAAAGGTGTGGTA
HC2E	-----
HC2F	-----
HC2A	GACACTCTTGAGCTGGACTTAGATTTTATTCTTCCTTGCAGAGTAGTGTTAGAATAGATG
HC2-80	GACACTCTTGAGCTGGACTTAGATTTTATTCTTCCTTGCAGAGTAGTGTTAGAATAGATG
HC2B	-----
HC2C	-----
HC2D-KIAA1058	GACACTCTTGAGCTGGACTTAGATTTTATTCTTCCTTGCAGAGTAGTGTTAGAATAGATG
HC2E	-----
HC2F	-----
HC2A	GCCTACAGAAAAAAAAGGTTCTGGGATCTACATGGCAGGGAGGGCTGCACTGACATTGAT
HC2-80	GCCTACAGAAAAAAAAGGTTCTGGGATCTACATGGCAGGGAGGGCTGCACTGACATTGAT
HC2B	-----
HC2C	-----
HC2D-KIAA1058	GCCTACAGAAAAAAAAGGTTCTGGGATCTACATGGCAGGGAGGGCTGCACTGACATTGAT
HC2E	-----
HC2F	-----
HC2A	GCCTGGGGGACCTTTTGCCTCGACTCGTGCCGGAAATCTGATCGTAATCAGGGTACAGAA
HC2-80	GCCTGGGGGACCTTTTGCCTCGACTCGTGCCGGAAATCTGATCGTAATCAGGGTACAGAA
HC2B	-----
HC2C	-----
HC2D-KIAA1058	GCCTGGGGGACCTTTTGCCTCGAGGCTGAGCTGGAAAATCTTGAAAATATTTTTT---T
HC2E	-----
HC2F	-----
HC2A	CTTACTAGTTTTGTCTAGGAGTATGTTGTATGACTAGGATTTGTGCTATTATCTCATTCA
HC2-80	CTTACTAGTTTTGTCTAGGAGTATGTTGTATGACTAGGATTTGTGCTATTATCTCATTCA
HC2B	-----
HC2C	-----
HC2D-KIAA1058	TTTCCTGTGGCACATTTCAGGTTGAATACAAGAACTATTTTTGTGACTAGTTTTTGATGAC
HC2E	-----
HC2F	-----
HC2A	ACAACATAGAGCAAGAATAGTGAGCTAACTGAGCTAGACACTCAATTAATCCGCTACTGG
HC2-80	ACAACATAGAGCAAGAATAGTGAGCTAACTGAGCTAGACACTCAATTAATCCGCTACTGG
HC2B	-----
HC2C	-----
HC2D-KIAA1058	CTAAGGGAACTGACCATTGTAATTTTTGTACCAGTGAACCAGGAGATTTAGTGCTTTTAT
HC2E	-----
HC2F	-----

FIG. 3A (cont.)

HC2A	CTTCAAGTCAGAACTTTGTCATTAATCATCGACTCCGGGACGGTCATATATGTATTACAT
HC2-80	CTTCAAGTCAGAACTTTGTCATTAATCATCGACTCCGGGACGGTCATATATGTATTACAT
HC2B	-----
HC2C	-----
HC2D-KIAA1058	ATTCATTTTCCTTGCATTTAAGAAAATATGAAAGCTTAAGGAATTATGTGAGCTTAAAACT
HC2E	-----
HC2F	-----
HC2A	TTCTACATTTTTTAATACTCACATGGGCTTATGCATTAAGTTTAATTGTGATAAATTTGTG
HC2-80	TTCTACATTTTTTAATACTCACATGGGCTTATGCATTAAGTTTAATTGTGATAAATTTGTG
HC2B	-----
HC2C	-----
HC2D-KIAA1058	AGTCAAGCAGTTTAGAACCAAAGGCCTATATTAATAACCGCAACTATGCTGAAAAGTACA
HC2E	-----
HC2F	-----
HC2A	CTGGTCCAGTATATGCAATACACTTTAATGGTTTATTCTTGTCATAAAAATGTGCAATAT
HC2-80	CTGGTCCAGTATATGCAATACACTTTAATGGTTTATTCTTGTCATAAAAATGTGCAATAT
HC2B	-----
HC2C	-----
HC2D-KIAA1058	AAGTAGTACAGTATATTGTTATGTACATATCATTGTTAATACAGTCCTGGCATTCTGTAC
HC2E	-----
HC2F	-----
HC2A	GGAGATGTATACAAGTCTTTACT-----
HC2-80	GGAGATGTATACAAGTCTTTACT-----
HC2B	-----
HC2C	-----
HC2D-KIAA1058	ATATATGTATTACATTTCTACATTTTTTAATACTCACATGGGCTTATGCATTAAGTTTAAT
HC2E	-----
HC2F	-----
HC2A	-----
HC2-80	-----
HC2B	-----
HC2C	-----
HC2D-KIAA1058	TGTGATAAATTTGTGCTGTTCCAGTATATGCAATACACTTTAATGTTTTATTCTTGTACA
HC2E	-----
HC2F	-----
HC2A	-----
HC2-80	-----
HC2B	-----
HC2C	-----
HC2D-KIAA1058	TAAAAATGTGCAATATGGAGATGTATACAGTCTTTACTATATTAGGTTTATAAACAGTTT
HC2E	-----
HC2F	-----

FIG. 3A (cont.)



HC2A	-----
HC2-80	-----
HC2B	-----
HC2C	-----
HC2D-KIAA1058	TAAGAATTTTCATCCTTTTGCCAAAATGGTGGAGTATGTAATTGGTAAATCATAAATCCTG
HC2E	-----
HC2F	-----

HC2A	-----
HC2-80	-----
HC2B	-----
HC2C	-----
HC2D-KIAA1058	TGGTGAATGGTGGTGTACTTTAAAGCTGTCACCATGTTATATTTTCTTTTAAGACATTAA
HC2E	-----
HC2F	-----

HC2A	-----
HC2-80	-----
HC2B	-----
HC2C	-----
HC2D-KIAA1058	TTTAGTAATTTTATATTTGGGAAAATAAAGGTTTTTAATTTTATTTAAGTGAATCACTG
HC2E	-----
HC2F	-----

HC2A	-----
HC2-80	-----
HC2B	-----
HC2C	-----
HC2D-KIAA1058	CCCTGCTGTAATTAAACATTCTGTACCACATCTGTATTAAAAAGACATTGCTGACC
HC2E	-----
HC2F	-----

FIG. 3A (cont.)

HC2A	-----
HC2A-80	-----
HC2B	-----
HC2C	-----
HC2D	ASGNLDKNARFSAIYRQDSNKLSNDDMLKLLADFRKPEKMAKLPVILGNLDITIDNVSSD
HC2E	-----
HC2F	-----

HC2A	-----
HC2A-80	-----
HC2B	-----
HC2C	-----
HC2D	FPNYVNSSYIPTKQFETCSKTPITFEVEEFVPCIPKHTQPYTIYTNHLYVYPKYLKYDSQ
HC2E	-----
HC2F	-----

HC2A	-----VLHHHQNPFFYDEIK
HC2A-80	-----
HC2B	-----
HC2C	-----
HC2D	KSFAKARNIAICIEFKDSDEEDSQPLKCIYGRPGGPVFTRSAFAAVLHHHQNPFFYDEIK
HC2E	-----
HC2F	-----

HC2A	IELPTQLHEKHHLTFFHVSCDNSSKGSTKKRDVVETQVGYSWLPLLKDGRVVTSEQHI
HC2A-80	-----
HC2B	-----
HC2C	-----
HC2D	IELPTQLHEKHHLTFFHVSCDNSSKGSTKKRDVVETQVGYSWLPLLKDGRVVTSEQHI
HC2E	-----
HC2F	-----

HC2A	PVSANLPSGYLGQELGMGRHYGPEIKWVDGGKPLLKISTHLVSTVYTQDQHLHNFFQYC
HC2A-80	-----
HC2B	-----
HC2C	-----
HC2D	PVSANLPSGYLGQELGMGRHYGPEIKWVDGGKPLLKISTHLVSTVYTQDQHLHNFFQYC
HC2E	-----
HC2F	-----

HC2A	QKTESGAQALGNELVKYLKSLHAMEGHVMIAFLPTILNQLFRVLTRATQEEVAVNVTRVI
HC2A-80	-----
HC2B	-----AMEGHVMIAFLPTILNQLFRVLTRATQEEVAVNVTRVI
HC2C	-----
HC2D	QKTESGAQALGNELVKYLKSLHAMEGHVMIAFLPTILNQLFRVLTRATQEEVAVNVTRVI
HC2E	-----AMEGHVMIAFLPTILNQLFRVLTRATQEEVAVNVTRVI
HC2F	-----

Fig. 3B

HC2A	IHVVAQCHEEGLESHLRSYVKYAYKAEPYVASEYKTVHEELTKSMTTILKPSADFLTSNK
HC2A-80	-----
HC2B	IHVVAQCHEEGLESHLRSYVKYAYKAEPYVASEYKTVHEELTKSMTTILKPSADFLTSNK
HC2C	-----
HC2D	IHVVAQCHEEGLESHLRSYVKYAYKAEPYVASEYKTVHEELTKSMTTILKPSADFLTSNK
HC2E	IHVVAQCHEEGLESHLRSYVKYAYKAEPYVASEYKTVHEELTKSMTTILKPSADFLTSNK
HC2F	-----

HC2A	LLRYSWFFFDVLIK SMAQH LIEN SKVKLLRNQRF PASYHHAAETVVNMLMPHITQKFGDN
HC2A-80	-----
HC2B	LLRYSWFFFDVLIK SMAQH LIEN SKVKLLRNQRF PASYHHAAETVVNMLMPHITQKFGDN
HC2C	-----
HC2D	LLKYSWFFFDVLIK SMAQH LIEN SKVKLLRNQRF PASYHHAAETVVNMLMPHITQKFRDN
HC2E	LLRYSWFFFDVLIK SMAQH LIEN SKVKLLRNQRF PASYHHAAETVVNMLMPHITQKFGDN
HC2F	-----

HC2A	PEASKNANHSLAVFIKRCFTFMDRGFVFKQINNYISCFAPGDPKTLFEYKFEFLRVVCNH
HC2A-80	-----
HC2B	PEASKNANHSLAVFIKRCFTFMDRGFVFKQINNYISCFAPGDPKTLFEYKFEFLRVVCNH
HC2C	-----
HC2D	PEASKNANHSLAVFIKRCFTFMDRGFVFKQINNYISCFAPGDPKTLFEYKFEFLRVVCNH
HC2E	PEASKNANHSLAVFIKRCFTFMDRGFVFKQINNYISCFAPGDPKTLFEYKFEFLRVVCNH
HC2F	-----

HC2A	EHYIPLNLPMPFGKGRIQRYQDLQLDYSLTDEF CRNHFLVGLLLREVG TALQEFREVRLI
HC2A-80	-----QLDYSLTDEF CRNHFLVGLLLREVG TALQEFREVRLI
HC2B	EHYIPLNLPMPFGKGRIQRYQDLQLDYSLTDEF CRNHFLVGLLLREVG TALQEFREVRLI
HC2C	-----
HC2D	EHYIPLNLPMPFGKGRIQRYQDLQLDYSLTDEF CRNHFLVGLLLREVG TALQEFREVRLI
HC2E	EHYIPLNLPMPFGKGRIQRYQDLQLDYSLTDEF CRNHFLVGLLLREVG TALQEFREVRLI
HC2F	-----

HC2A	AISVLKNLLIKHSFDDRYASRSHQARIATLYLPLFGLLIENVQRINVRDVSPFPVNAGMT
HC2A-80	AISVLKNLLIKHSFDDRYASRSHQARIATLYLPLFGLLIENVQRINVRDVSPFPVNAGMT
HC2B	AISVLKNLLIKHSFDDRYASRSHQARIATLYLPLFGLLIENVQRINVRDVSPFPVNAGMT
HC2C	-----
HC2D	AISVLKNLLIKHSFDDRYASRSHQARIATLYLPLFGLLIENVQRINVRDVSPFPVNAGMT
HC2E	AISVLKNLLIKHSFDDRYASRSHQARIATLYLPLFGLLIENVQRINVRDVSPFPVNAGMT
HC2F	-----

HC2A	VKDESLALPAVNPLVTPQKGSTLDNSLHKDLLGAISGIASPYTTSTPNINSVRNADSRGS
HC2A-80	VKDESLALPAVNPLVTPQKGSTLDNSLHKDLLGAISGIASPYTTSTPNINSVRNADSRGS
HC2B	VKDESLALPAVNPLVTPQKGSTLDNSLHKDLLGAISGIASPYTTSTPNINSVRNADSRGS
HC2C	-----
HC2D	VKDESLALPAVNPLVTPQKGSTLDNSLHKDLLGAISGIASPYTTSTPNINSVRNADSRGS
HC2E	VKDESLALPAVNPLVTPQKGSTLDNSLHKDLLGAISGIASPYTTSTPNINSVRNADSRGS
HC2F	-----ADSRGS

Fig. 3B (cont.)

HC2A	LISTDSGNSLPERNSEKSNSLDKHQQSSTLGNSVVRCDKLDQSEIKSLLMCFLYILKSMS
HC2A-80	LISTDSGNSLPERNSEKSNSLDKHQQSSTLGNSVVRCDKLDQSEIKSLLMCFLYILKSMS
HC2B	LISTDSGNSLPERNSEKSNSLDKHQQSSTLGNSVVRCDKLDQSEIKSLLMCFLYILKSMS
HC2C	-----
HC2D	LISTDSGNSLPERNSEKSNSLDKHQQSSTLGNSVVRCDKLDQSEIKSLLMCFLYILKSMS
HC2E	LISTDSGNSLPERNSEKSNSLDKHQQSSTLGNSVVRCDKLDQSEIKSLLMCFLYILKSMS
HC2F	LISTDSGNSLPERNSEKSNSLDKHQQSSTLGNSVVRCDKLDQSEIKSLLMCFLYILKSMS
HC2A	DDALFTYWNKASTSELMDFFTISEVCLHQFQYMGKRYIARNQEGLGPIVHDRKSQTLPV
HC2A-80	DDALFTYWNKASTSELMDFFTISEVCLHQFQYMGKRYIARNQEGLGPIVHDRKSQTLPV
HC2B	DDALFTYWNKASTSELMDFFTISEVCLHQFQYMGKRYIARNQEGLGPIVHDRKSQTLPV
HC2C	-----
HC2D	DDALFTYWNKASTSELMDFFTISEVCLHQFQYMGKRYIAR-----
HC2E	DDALFTYWNKASTSELMDFFTISEVCLHQFQYMGKRYIARNQEGLGPIVHDRKSQTLPV
HC2F	DDALFTYWNKASTSELMDFFTISEVCLHQFQYMGKRYIAS-----VR--KISSVLGIS
HC2A	RNRTGMMHARLQQLGSLDNSLTFNHSYGHSDADVLHQSLLEANIATEVCLTALDTLSLFT
HC2A-80	RNRTGMMHARLQQLGSLDNSLTFNHSYGHSDADVLHQSLLEANIATEVCLTALDTLSLFT
HC2B	RNRTGMMHARLQQLGSLDNSLTFNHSYGHSDADVLHQSLLEANIATEVCLTALDTLSLFT
HC2C	-----
HC2D	---TGMHARLQQLGSLDNSLTFNHSYGHSDADVLHQSLLEANIATEVCLTALDTLSLFT
HC2E	RNRTGMMHARLQQLGSLDNSLTFNHSYGHSDADVLHQSLLEANIATEVCLTALDTLSLFT
HC2F	V-----D-NG-----YGHSDADVLHQSLLEANIATEVCLTALDTLSLFT
HC2A	LAFKNQLLADHGHNPLMKKVFDVYLCFLQKHQSETALKNVFTALRS LIYKFPSTFYEGRA
HC2A-80	LAFKNQLLADHGHNPLMKKVFDVYLCFLQKHQSETALKNVFTALRS LIYKFPSTFYEGRA
HC2B	LAFK--LLADHGHNPLMKKVFDVYLCFLQKHQSETALKNVFTALRS LIYKFPSTFYEGRA
HC2C	-----
HC2D	LAFKNQLLADHGHNPLMKKVFDVYLCFLQKHQSETALKNVFTALRS LIYKFPSTFYEGRA
HC2E	LAFKNQLLADHGHNPLMKKVFDVYLCFLQKHQSETALKNVFTALRS LIYKFPSTFYEGRA
HC2F	LAFKNQLLADHGHNPLMKKK-----
HC2A	DMCAALCYEILKCCNSKLSSIRTEASQLLYFLMRNNFDYTGKKS FVRTHLQV IISVSQ LI
HC2A-80	DMCAALCYEILKCCNSKLSSIRTEASQLLYFLMRNNFDYTGKKS FVRTHLQV IISVSQ LI
HC2B	DMCAALCYEILKCCNSKLSSIRTEASQLLYFLMRNNFDYTGKKS FVRTHLQV IISVSQ LI
HC2C	-----
HC2D	DMCAALCYEILKCCNSKLSSIRTEASQLLYFLMRNNFDYTGKKS FVRTHLQV IISVSQ LI
HC2E	DMCAALCYEILKCCNSKLSSIRTEASQLLYFLMRNNFDYTGKKS FVRTHLQV IISVSQ LI
HC2F	-----
HC2A	ADVVGIGETR FQQSLS I INNCANS DR LIKHTS FSSDV KDLTKR IRTVLMATAQMKEHEND
HC2A-80	ADVVGIGETR FQQSLS I INNCANS DR LIKHTS FSSDV KDLTKR IRTVLMATAQMKEHEND
HC2B	ADVVGIGETR FQQSLS I INNCANS DR LIKHTS FSSDV KDLTKR IRTVLMATAQMKEHEND
HC2C	-----
HC2D	ADVVGIGETR FQQSLS I INNCANS DR LIKHTS FSSDV KDLTKR IRTVLMATAQMKEHEND
HC2E	ADVVGIGETR FQQSLS I INNCANS DR LIKHTS FSSDV KDLTKR IRTVLMATAQMKEHEND
HC2F	-----

FIG. 3B (cont.)

HC2A	PEMLVDLQYSLAKSYASTPELRKTWLD SMARIHVKN GDLSEAAMCYVHV TALVAEYLTRK
HC2A-80	PEMLVDLQYSLAKSYASTPELRKTWLD SMARIHVKN GDLSEAAMCYVHV TALVAEYLTRK
HC2B	PEMLVDLQYSLAKSYASTPELRKTWLD SMARIHVKN GDLSEAAMCYVHV TALVAEYLTRK
HC2C	-----
HC2D	PEMLVDLQYSLAKSYASTPELRKTWLD SMARIHVKN GDLSEAAMCYVHV TALVAEYLTRK
HC2E	PEMLVDLQYSLAKSYASTPELRKTWLD SMARIHVKN GDLSEAAMCYVHV TALVAEYLTRK
HC2F	-----

HC2A	-----GVFRQGCTAFRVITPNIDEEASMMEDVGMQDVHFNE
HC2A-80	-----GVFRQGCTAFRVITPNIDEEASMMEDVGMQDVHFNE
HC2B	-----GVFRQGCTAFRVITPNIDEEASMMEDVGMQDVHFNE
HC2C	-----FRQGCTAFRVITPNIDEEASMMEDVGMQDVHFNE
HC2D	EAVQWEPPLLPHSHSACLRRSRGGVFRQGCTAFRVITPNIDEEASMMEDVGMQDVHFNE
HC2E	-----GVFRQGCTAFRVITPNIDEEASMMEDVG-----
HC2F	-----

HC2A	DVLMELLEQCADGLWKAERYELIADIYKLIIP IYEKRR-----
HC2A-80	DVLMELLEQCADGLWKAERYELIADIYKLIIP IYEKRR-----
HC2B	DVLMELLEQCADGLWKAERYELIADIYKLIIP IYEKRRDFERLAHLYDTLHRAYSK
HC2C	DVLMELLEQCADGLWKAERYELIADIYKLIIP IYEKRRDFERLAHLYDTLHRAYSK
HC2D	DVLMELLEQCADGLWKAERYELIADIYKLIIP IYEKRRDFERLAHLYDTLHRAYSK
HC2E	-----KAERYELIADIYKLIIP IYEKRRDFERLAHLYDTLHRAYSK
HC2F	-----

HC2A	-----DFFEDEDGKEYIYKEPKLTPLSE
HC2A-80	-----DFFEDEDGKEYIYKEPKLTPLSE
HC2B	VTEVMHSGRLLGTYFRVAFFGQ-----GFFEDEDGKEYIYKEPKLTPLSE
HC2C	VTEVMHSGRLLGTYFRVAFFGQ-----GFFEDEDGKEYIYKEPKLTPLSE
HC2D	VTEVMHSGRLLGTYFRVAFFGQAAQYQFTDSETDVEGFFEDEDGKEYIYKEPKLTPLSE
HC2E	VTEVMHSGRLLGTYFRVAFFGQ-----GFFEDEDGKEYIYKEPKLTPLSE
HC2F	-----

HC2A	ISQRLCLKLYSDKFGSENVKMIQDSGKVNPKDLDSKYAYIQVTHVIPFFDEKELQERKTEF
HC2A-80	ISQRLCLKLYSDKFGSENVKMIQDSGKVNPKDLDSKYAYIQVTHVIPFFDEKELQERKTEF
HC2B	ISQRLCLKLYSDKFGSENVKMIQDSGKVNPKDLDSKYAYIQVTHVIPFFDEKELQERKTEF
HC2C	ISQRLCLKLYSDKFGSENVKMTQDSGKVNPKDLDSKYAYIQVTHVIPFFDEKELQERKTEF
HC2D	ISQRLCLKLYSDKFGSENVKMIQDSGKVNPKDLDSKYAYIQVTHVIPFFDEKELQERKTEF
HC2E	ISQRLCLKLYSDKFGSENVKMIQDSGKVNPKDLDSKYAYIQVTHVIPFFDEKELQERKTEF
HC2F	-----

HC2A	ERSHNIRRFMFEMPFTQTGKRQGGVEEQCKRRTILTAIHCFPYVKKRIPV MYQHHTDLNP
HC2A-80	ERSHNIRRFMFEMPFTQTGKRQGGVEEQCKRRTILTAIHCFPYVKKRIPV MYQHHTDLNP
HC2B	ERSHNIRRFMFEMPFTQTGKRQGGVEEQCKRRTILTAIHCFPYVKKRIPV MYQHHTDLNP
HC2C	ERSHNIRRFMFEMPFTQTGKRQGGVEEQCKRRTILTAIHCFPYVKKRIPV MYQHHTDLNP
HC2D	ERSHNIRRFMFEMPFTQTGKRQGGVEEQCKRRTILTAIHCFPYVKKRIPV MYQHHTDLNP
HC2E	ERSHNIRRFMFEMPFTQTGKRQGGVEEQCKRRTILTAIHCFPYVKKRIPV MYQHHTDLNP
HC2F	-----

FIG. 3B (cont.)



HC2A  
HC2A-80  
HC2B  
HC2C  
HC2D  
HC2E  
HC2F

IEVAIDEMSKKVAELRQLCSSAEVDMIKLQLKLQGSVSVQVNAGPLAYARAFLLDDTNTKR  
IEVAIDEMSKKVAELRQLCSSAEVDMIKLQLKLQGSVSVQVNAGPLAYARAFLLDDTNTKR  
IEVAIDEMSKKVAELRQLCSSAEVDMIKLQLKLQGSVSVQVNAGPLAYARAFLLDDTNTKR  
IEVHZ-----  
IEVAIDEMSKKVAELRQLCSSAEVDMIKLQLKLQGSVSVQVNAGPLAYARAFLLDDTNTKR  
IEVAIDEMSKKVAELRQLCSSAEVDMIKLQLKLQGSVSVQVNAGPLAYARAFLLDDTNTKR  
-----

HC2A  
HC2A-80  
HC2B  
HC2C  
HC2D  
HC2E  
HC2F

YPDNKVKLLKEVFRQFVEACGOALAVNERLIKEDQLEYQEEMKANYREMAKELSEIMHEQ  
YPDNKVKLLKEVFRQFVEACGOALAVNERLIKEDQLEYQEEMKANYREMAKELSEIMHEQ  
YPDNKVKLLKEVFRQFVEACGOALAVNERLIKEDQLEYQEEMKANYREMAKELSEIMHEQ  
-----  
YPDNKVKLLKEVFRQFVEACGOALAVNERLIKEDQLEYQEEMKANYREMAKELSEIMHEQ  
YPDNKVKLLKEVFRQFVEACGOALAVNERLIKEDQLEYQEEMKANYREMAKELSEIMHEQ  
-----

HC2A  
HC2A-80  
HC2B  
HC2C  
HC2D  
HC2E  
HC2F

ICPLEEKTSVLPNSLHIFNAISGTPTSTMVHGMTSSSSVVZ----  
ICPLEEKTSVLPNSLHIFNAISGTPTSTMVHGMTSSSSVVZ----  
ICPLEEKTSVLPNSLHIFNAISGTPTSTMVHGMTSSSSVVZ----  
-----  
LG-----  
ICPLEEKTSVLPNSLHIFNAISGTPTSTMVHGMTSSSSVVZ----  
-----

FIG. 3B(cont.)

006707 FEB 85

PBL  
lung  
placenta  
sm intestine  
liver  
kidney  
spleen  
thymus  
colon  
skel muscle  
heart  
brain



FIG. 4A

00ETOT" /EB/8960

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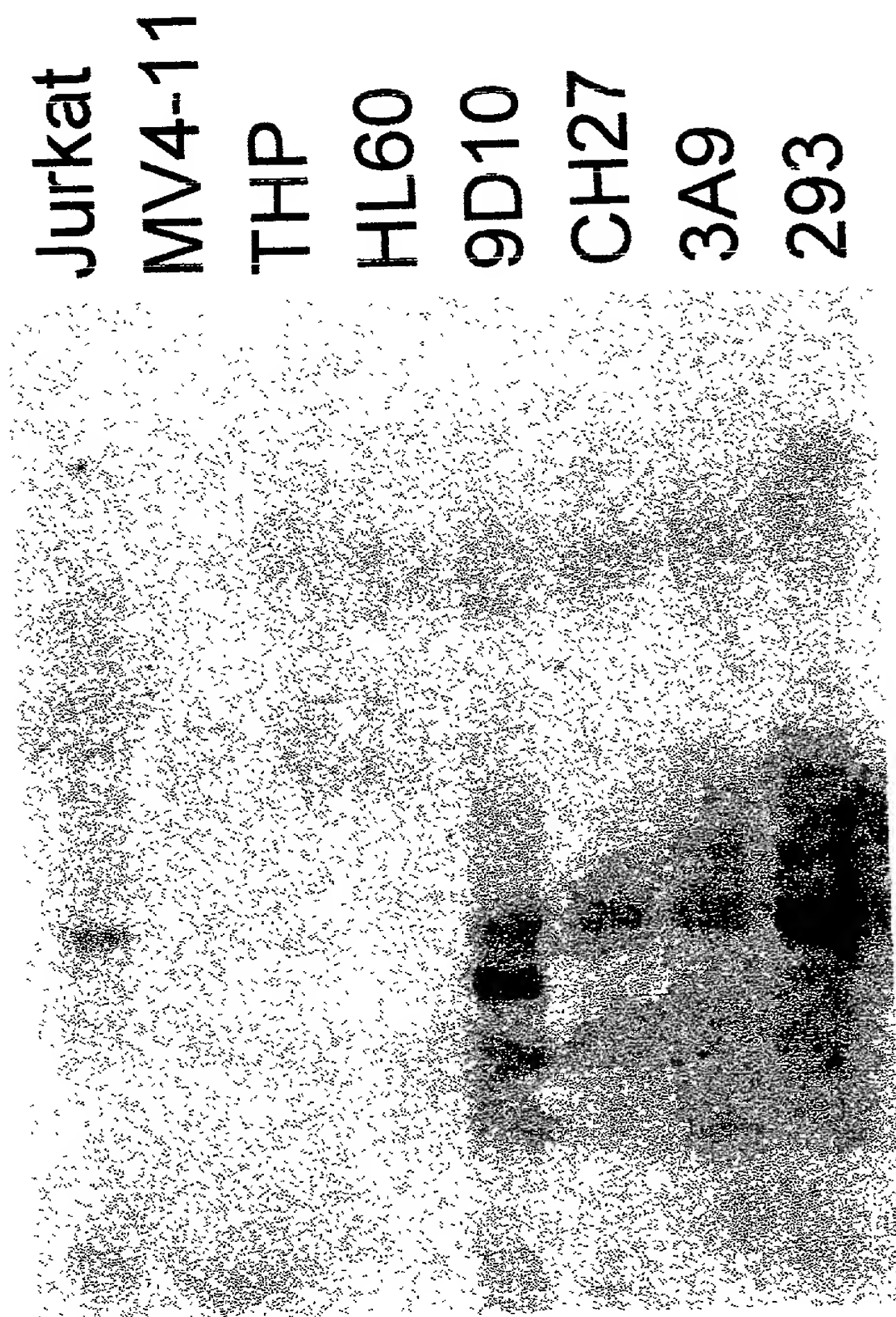


Fig. 4B

HC2A	-----
KIAA	ASGNLDKNARFSAIYRQDSNKLSNDDMLKLLADFRKPEKMAKLPVILGNLDITIDNVSSD
rat	-----
HC4	-----
HC1	-----
HC3	-----
HC5	-----

HC2A	-----
KIAA	FPNYVNSSYIPTKQFETCSKTPITFEVEEFVPCIPKHTQPYTIYTNHLYVYPKYLKYDSQ
rat	-----
HC4	-----
HC1	-----
HC3	-----
HC5	-----

HC2A	-----VLHHHQNPETYDEIK
KIAA	KSFAKARNIAICIEFKDSDEEDSQPLKCIYGRPGGPVFTRSAFAAVLHHHQNPETYDEIK
rat	-----
HC4	-----
HC1	-----
HC3	-----
HC5	-----

HC2A	IELPTQLHEKHLLLLTFFHVSCDNSSKGSTKKRDVVETQVGYSWLPLLKDGRVVTSEQHI
KIAA	IELPTQLHEKHLLLLTFFHVSCDNSSKGSTKKRDVVETQVGYSWLPLLKDGRVVTSEQHI
rat	-----
HC4	-----
HC1	-----
HC3	-----
HC5	-----

HC2A	PVSANLPSGYLGYQELGMGRHYGPEIKWVDGGKPLLKISTHLVSTVYTQDQHLHNFFQYC
KIAA	PVSANLPSGYLGYQELGMGRHYGPEIKWVDGGKPLLKISTHLVSTVYTQDQHLHNFFQYC
rat	-----
HC4	-----
HC1	-----
HC3	-----GPGPARSTVSISLISNSARV
HC5	-----

HC2A	QKTESGAQALGNELVKYLKSLHAMEGHVMIAFLPTILNQLFRVLT-RATQEEVAVNVTRV
KIAA	QKTESGAQALGNELVKYLKSLHAMEGHVMIAFLPTILNQLFRVLT-RATQEEVAVNVTRV
rat	-----
HC4	-----MEIQVLIRFLSVILMQLEFWVLPNMIHEDDVPISCPMPV
HC1	-----MSFLPIILNQLFKVLV-QNEEDEITTTVTRV
HC3	NRSRSLSNSNPDISGTPTSPDDEVRSIIGSKGLDRSNSWVNTGGPKAAPWGSNPSPSAES
HC5	-----

Fig. 5A

HC2A	I I H V V A Q C H E E G L E S H L R S Y V K Y A Y K A E P Y V A S E Y K T V H E E L T K S M T T I L K P S A D F L T S N
KIAA	I I H V V A Q C H E E G L E S H L R S Y V K Y A Y K A E P Y V A S E Y K T V H E E L T K S M T T I L K P S A D F L T S N
rat	-----
HC4	L F H I V S K C H E E G L D S Y L S S F I K Y S F R P G K P S A P Q A P L I H E T L A T M M I A L L K Q S A D F L A I N
HC1	L P D I V A K C H E E Q L D H S V Q S Y I K F V F K T R --- A C K E R P V H E D L A K N V T G L L K - S N D S P T V K
HC3	T Q A M D R S C N R M S S H T E T S S F L Q T L T G R L P --- T K K L F H E E L A L Q W V V C S G -- S V R --- E
HC5	-----

Cadherin  
Cleavage

HC2A	K L L R Y S W F F F D V L I K S M A Q H L I E N S K V K L L R N Q R F P A S Y H H A A E T V V N M L M P H I T Q K F G D
KIAA	K L L K Y S W F F F D V L I K S M A Q H L I E N S K V K L L R N Q R F P A S Y H H A V E T V V N M L M P H I T Q K F R D
rat	-----
HC4	K L L K Y S W F F F E I I A K S M A T Y L L E E N K I K L T H G Q R F P K A Y H H A L H S L F L A I T - I V E S Q Y A E
HC1	H V L K H S W F F F A I I L K S M A Q H L I D T N K I Q L P R P Q R F P E S Y Q N E L D N L V M V L S D H V I W K Y K D
HC3	S A L Q Q A W F F F E L M V K S M V H H L Y F N D K L E A P R K S R F P E R F M D D I A A L V S T I A S D I V S R F Q K
HC5	-----

HC2A	N P E A S K N A N H S L A V F I K R C F T F M D R G F V F K Q I N --- N Y I S -- C F A P G D P K T L F E Y K F E F L
KIAA	N P E A S K N A N H S L A V F I K R C F T F M D R G F V F K Q I N --- N Y I S -- C F A P G D P K T L F E Y K F E F L
rat	-----
HC4	I P K E S R N V N Y S L A S F L K C C L T L M D R G F V F N L I N --- D Y I S -- G F S P K D P K V L A E Y K F E F L
HC1	A L E E T R R A T H S V A R F L K R C F T F M D R G C V F K M V N --- N Y I S -- M F S S G D L K T L C Q Y K F D F L
HC3	D T E M V E R L N T S L A F F L N D L L S V M D R G F V F S L I K S C Y Q V S S K L Y S L P N P S V L V S L R L D F L
HC5	-----

HC2A	R V V C N H E H Y I P L N L P M --- P F G K G R I Q R --- Y Q D L Q L --- D Y S L T D E F
KIAA	R V V C N H E H Y I P L N L P M --- P F G K G R I Q R --- Y Q D L Q L --- D Y S L T D E F
rat	-----
HC4	Q T I C N H E H Y I P L N L P M --- A F A K P K L Q R --- V Q D S N L --- E Y S L S D E Y
HC1	Q E V C Q H E H F I P L C L P I R S A N I P D P L T P S E S --- T Q E L H A S D M P E Y S V T N E F
HC3	R I I C S H E H Y V T L N L P C S L L T P P A S P S P S V S S A T S Q S S G F S T N V Q D Q K I A N M F E L S -- V P F
HC5	----- M N A D T A P T S P C P S I S --- S Q N S S C S S F Q D Q K I A S M F D R T S R V P A

HC2A	C R N H F L V G L L L R E V G T A L Q E F R E --- V R L I A I S V L K N L L I K H S F D D R Y A S R S H Q A R I A T
KIAA	C R N H F L V G L L L R E V G T A L Q E F R E --- V R L I A I S V L K N L L I K H S F D D R Y A S R S H Q A R I A T
rat	-----
HC4	C K H H F L V G L L L R E T S I A L Q D N Y E --- I R Y T A I S V I K N L L I K H A F D T R Y Q H K N Q Q A K I A Q
HC1	C R K H F L I G I L L R E V G F A L Q E D Q D --- V R H L A L A V L K N L M A K H S F D D R Y R E P R K Q A Q I A S
HC3	R Q Q H Y L A G L V L T E L A V I L D P D A E G L F G L H K K V I N M V H N L L S S H D S D P R Y S D P Q I K A R V A M
HC5	S S T S - S P G L L F T E L A A A L D A E G E G I S E V Q R K A V S A I H S L L S S H D L D P R C V K P E V K V K I A A

HC2A	L Y L P L F G L L I E N V Q R I N V R D V S P F P V N A G - M T V K D E S L A L P A V N P L V T P Q K G S T L D N S L H
KIAA	L Y L P L F G L L I E N V Q R I N V R D V S P F P V N A G - M T V K D E S L A L P A V N P L V T P Q K G S T L D N S L H
rat	-----
HC4	L Y L P F V G L L L E N I Q R L A G R D T L Y S C A A M P N S A S R D E F P C G --- F T S P -- A N -- R G S L S
HC1	L Y M P L Y G M L L D N M P R I Y L K D L Y P F T V N T S N Q G S R D D L S T N G G F Q S Q T A I K H A N S V D T S F S
HC3	L Y L P L I G I I M E T V P Q L Y D F T E T H N Q R G R P I C I A T D D Y E S E --- S G --- S M I S
HC5	L Y L P L V G I I L D A L P Q L C D F T V A D T R R Y R --- T S G S D E E Q E --- G A --- G A I T

HC2A	K D L L G A I S G I A S P Y T T S T P N I N S V R N A D S R G S L I S T D S G N S L P E R N S E K S N S L D K H Q Q S S
KIAA	K D L L G A I S G I A S P Y T T S T P N I N S V R N A D S R G S L I S T D S G N S L P E R N S E K S N S L D K H Q Q S S
rat	-----
HC4	T D K D T A Y G S F Q N G --- H G I K R E D S R G S L I P - E G A T G F P D Q G N T G E N --- T R Q S
HC1	K D V L N S I A A F S S --- I A I S T V N H A D S R A S L A S L D S N P S T N E K S S E K T D N C E K I P R P L
HC3	Q T V A M A I A G T S V P Q --- L T R P G S F L L T S T S G R Q H T ---
HC5	Q N V A L A I A G N N F N --- L K T S G - I V L S S L P Y K Q Y N ---



HC2A	TLGNSVVRCDKLDQSEIKSLLMCFLYILKSMSDDALFTYWN-KASTSELMDFFTISEVCL
KIAA	TLGNSVVRCDKLDQSEIKSLLMCFLYILKSMSDDALFTYWN-KASTSELMDFFTISEVCL
rat	-----
HC4	STRSSVSQYNRLDQYEIRSLLMCYLYIVKMISEDTLTYWN-KVSPQELINILILLEVCL
HC1	ALIGSTLRFDRDLQAETRSLLMCFLHIMKTISYETLIAYWQ-RAPSPEVSDFFSILDVCL
HC3	-----TFSAESSRSLICLLWVLKN-ADETVLQKWFTDLSVLQLNRLDLLYLCV
HC5	-----MLNADTTRNLMICFLWIMKN-ADQSLIRKWIADLPSTQLNRILDLLFICV

HC2A	HQFQYMGKRYIARNQEGLG--PIVHDRKS-----QTLPVSRNRTGMM
KIAA	HQFQYMGKRYIAR-----TGMM
rat	-----
HC4	FHFRYMGKRNIARVHDAWLSKHFGIDRKS-----QTMPALRNRSGVM
HC1	QNFYRLGKRNIIRKIAAAF--KFVQSTQNNGLKGSNPSCQTSGLLAQWMHSTSRHEGHK
HC3	SCFEYKGGKVFERNLSLTFK--KSKDMRAK-----LEEAILGSIGARQEMV
HC5	LCFEYKGGKQSSDKVSTQVLQ--KSRDVkar-----LEEALLRGEgARGEMM

HC2A	HARLQQL-----GSLDNS-----LTFNHSYGHSDADVLHQSLLEANIATEVC
KIAA	HARLQQL-----GSLDNS-----LTFNHSYGHSDADVLHQSLLEANIATEVC
rat	-----
HC4	QARLQHL-----SSLESS-----FTLNHSSTTTEADI FHQALLEGNTATEVS
HC1	QHRSTLPIIRGK--NALSnpKL---LQMLDNTMTSnsNEIDIVHHVDTEANIATEGC
HC3	RRSRGQLERSPSGSafGSQENLRWRKDMTHWRQNTekLDKSRAEIEHEALIDGNLATEAN
HC5	RRRAPGNDRFP-----GLNENLRWKKEQTHWRQANEKLDKTKAELDQeALISGNLATEAH

HC2A	LTALDTLSLFTLAFKNQLLADHGHNPIMKKVFDVYLCFLQKHQSETALKNVFTALRSliY
KIAA	LTALDTLSLFTLAFKNQLLADHGHNPIMKKVFDVYLCFLQKHQSETALKNVFTALRSliY
rat	-----KLSRGHSPLMKKVFdvYLCFLQKHQSEMALKNVFTALRSliY
HC4	LTVLDTISFFTQCFKTHFLNNDGHNPIMKKVFDIHLAFLKNGQSEVSLKHVFASLRAFIS
HC1	LTILDVLSLFTQTHQRQLQQCDQNSLMKRGFDTYMLFFQVNQSATALKHVFASLRLFVC
HC3	LIILDTLEIVVQTVS--VTES--KESILGGVLKvLLHSMACNQSAVYLQHCfATQRALVS
HC5	LIILDMQENIIQASS--ALDC--KDSLLGGVLRVLVNSLNCdQSTTYLTHCFATLRAlIA

HC2A	KFPSTFYEGRADMCALCYEILKCCNSKLSSIRTEASQLLYFLMRNnFDYTgKKSfVRTH
KIAA	KFPSTFYEGRADMCALCYEILKCCNSKLSSIRTEASQLLYFLMRNnFDYTgKKSfVRTH
rat	KFPSTFYEGRADMCASLCYEVlKCCNSKLSSIRTEASQLLYFLMRNnFDYTgKKSfVRTH
HC4	KFPSAFFKGRVNMCAAFCYEVlKCCTSKISTRNEASALLYLLMRNnFEYTKRKTFLRTH
HC1	KFPSAFFQGPADLCGSFCYEVlKCCNHRSRSTQTEASALLYLFMRKNFEFNKQKSIVRSH
HC3	KFPELLFEEETEQCADLCLRLLRHCSSSIGTIRSHPSASLYLLMRQNFElGN--NFARVK
HC5	KFGDLLFEEEEVEQCFDLCHQVLHHCSSSMDVTRSQACATLYLLMRFSFGATS--NFARVK

HC2A	LQVIISVSQLIADVVGIGETRfQQSLSIINNCANSdRLIKHTSFSSDVkdLTkRIRTVLM
KIAA	LQVIISVSQLIADVVGIGETRfQQSLSIINNCANSdRLIKHTSFSSDVkdLTkRIRTVLM
rat	LQVIISLSQLIADVVGIGETRfQQSLSIINNCANSdRLIKHTSFSSDVkdLTkRIRTVLM
HC4	LQIIIAVSQLIADVALSGGSrfQESLFIINNFANSdRPMlARAFPAEVkdLTkRIRTVLM
HC1	LQLIKAVSQLIAD-AGIGGSrfQHSLAITNnFANGDKQMKNsNFPAEVkdLTkRIRTVLM
HC3	MQVMSLSSLVGTSQNFNEEFLLRSLKTIltYAEEDLELRETTFPDQVQDLVFNlHMILS
HC5	MQVTMSLASLVGRAPDFNEEHLRSLRTIlAYSEEDTAMQMTpFPtQVEELLcNLNSIlY

FIG. 5A (cont.)

HC2A	ATAQMKEHENDPEMLVDLQYSLAKSYASTPELRKTWLD	SMARIHVKN	GD	LSEAAMCYVHV
KIAA	ATAQMKEHENDPEMLVDLQYSLAKSYASTPELRKTWLD	SMARIHVKN	GD	LSEAAMCYVHV
rat	ATAQMKEHENDPEMLVDLQYSLAKSYASTPELRKTWLD	SMARIHVKN	GD	LSEAAMCYVHV
HC4	ATAQMKEHEKDPEMLIDLQYSLAKSYASTPELRKTWLD	MAKIHVKN	GD	FSEAAMCYVHV
HC1	ATAQMKEHEKDPEMLVDLQYSLANSYASTPELRRTWLES	MAKIHARNG	DL	LSEAAMCYIHI
HC3	DTVKMKEHQEDPEMLIDLMYRIAKGYQTSPDLRLTLWLQ	NMAGKHSERSN	HA	EAAQCLVHS
HC5	DTVKMREFQEDPEMLMDLMYRIAKSYQASPDRLTLWLQ	NMAEKHTKKK	CY	TEAMCLVHA

## SH3

HC2A	TALVAEYI	TRKGV	---	---	FRQGCTAFRVITPN
KIAA	TALVAEYI	TRKEA	---	---	VQWEPPLLPHSHSACLRRSRGGVFRQGCTAFRVITPN
rat	TALVAEYI	TRKEAD	---	---	LALQREPPVFPYSHTSCQKSRGGMFRQGCTAFRVITPN
HC4	AALVAEFI	HRKKL	---	---	FPNGCSAFKKITPN
HC1	AALIAEYI	KRKGWYKVEKIC	IASLLSE	THPCDSNSLLTTPSGGSMFSGMGP	AFLSITPN
HC3	AALVAEYI	SMLD	---	---	RKYLPGCVTFQNISSN
HC5	AALVAEYI	SMLD	---	---	HSYLPVGSVSFQNISSN

HC2A	IDEEASMMEDVGMQD	---	---	VHFNEDVLMELLEQCADGLWKAERYELIADIYKLIIP
KIAA	IDEEASMMEDVGMQD	---	---	VHFNEDVLMELLEQCADGLWKAERYELIADIYKLIIP
rat	IDEEASMMEDVGMQD	---	---	VHFNEDVLMELLEQCADGLWKAERLRAGLLTSINSSSP
HC4	IDEEGAMKEDAGMMD	---	---	VHYSEEVLLLELLEQCVNGLWKAERYEIISEISKLIGPI
HC1	IKEEGAAKEDSGMHD	---	---	TPYNENILVEQLYMCGEFLWKSEYELIADVNPPIIAV
HC3	VLEESAVSDDVSPDEEGICSGKYFTESGLVGLLEQAAASF	SMAGMYEAVNEVYKVLIP		
HC5	VLEESVVS	EDTLSPDE	DGVCAGQYFTESGLVGLLEQAAELFSTGGLYETVNEVYKVLIP	

		ITAM	ITAM		ITAM	ITAM
HC2A	YEKRRD	---	---	---	---	---
KIAA	YEKRRD	FERLAHLYDTLHRA	YSKVTEVMHSGRRL	LGTYFRVAFFGQAAQYQFTDSETDVE		
rat	SMKSGGTLETTHLYDTLH	RPYSKVTEVITR	---	---	A---	AGSWDLLPGGLFGQ
HC4	YENRREFENLTQVYRTLHG	AYTKILEVMHTKKRLLG	---	---	---	TFFRVAFYGQ
HC1	FEKQRDFKKLSDLIYYDIH	RSYLKVAEVVNSEKRLFG	---	---	---	RYFRVAFYGQ
HC3	HEANRDAKKLSTIHGKLQ	EAFSKI	VHQTGWERMFG	---	---	TYFRVGFYG-
HC5	LEAHREFRKLTLTHSKLQ	RAFD	SVNKH--KRMFG	---	---	TYFRVGFFG-

		ITAM		ITAM	
HC2A	-FFED	EDGKEYIYKEPKLTPLSEISQRL	LLKLYSDK	FGSENVKMIQDSGKVNPKDLDSKYA	
KIAA	GFFE	EDGKEYIYKEPKLTPLSEISQRL	LLKLYSDK	FGSENVKMIQDSGKVNPKDLDSKYA	
rat	GFFE	EDGKEYIYKEPKLTPLSEISQRL	LLKLYSDK	FGSENVKMIQDSGKVNPKDLDSKFA	
HC4	SFFE	EEDGKEYIYKEPKLTGLSEISLRLV	KLYGEKFGTENVKIIQSDKVN	AKELDPKYA	
HC1	GFFE	EEEGKEYIYKEPKLTGLSEISQRL	LLKLYADKFGADNVKIIQDSN	KVNPKDLDPKYA	
HC3	TKFGDLDEQEFVYKEFAIT	KLAEISHRLEGEFYGERFGEDVVEVIKDSNPVDKCKLDPNKA			
HC5	SKFGDLDEQEFVYKEFAIT	KLPEISHRLEAFYGO	CFGAEFVEVIKdstpvdktklDPNKA		

## ITAM

HC2A	YIQVTHVIPFFDEKELQERKTEFERSHNIRRFMFEMPFTQTGKRQGGVVEEQCKRRTILTA
KIAA	YIQVTHVIPFFDEKELQERKTEFERSHNIRRFMFEMPFTQTGKRQGGVVEEQCKRRTILTA
rat	YIQVTHVTPFFDEKELQERKTEFERCHNIRRFMFEMPFTQTGKRQGGVVEEQCKRRTILTA
HC4	HIQVTYVVKPYFDDKELTERKTEFERNHNISRFVFEAPYTLGKKGQCIIEEQCKRRTILT
HC1	YIQVTYVTPFFEEKEIEDRKTDFEMHHNINRFVFETPFTLSGKKHGGVAEQCKRRTILT
HC3	YIQITYVEPYFDYEMKDRITYFDKNYNLRREMYCTPFTLDGRAHGEHQFQRKTILT
HC5	YIQITFVEPYFDEYEMKDRVITYFEKNFNLRREMYTTPFTLEGRPRGELHEQYRRNTVLT

FIG. 5A (cont.)

## Coiled-Coil 1

HC2A	IHC	FPYVKKRIPVMYQHHTD	LNPIEVAIDEMSKKVAELRQLCSSAEVDMIKLQ	KLQGSV
KIAA	IHC	FPYVKKRIPVMYQHHTD	LNPIEVAIDEMSKKVAELRQLCSSAEVDMIKLQ	KLQGSV
rat	IHC	FPYVKKRIPVMYQHHTD	LNPIEVAIDEMSKKVAELHQLCSSAEVDMIKLQ	KLQGSV
HC4	SNS	FPYVKKRIPINCEQQINIKPIDGATDEIKDKTAE	LQKLCSSSTDVDMIQLQ	KLQGWV
HC1	SHL	FPYVKKRIQVISQSSTE	LNPIEVAIDEMSRKVSELNQLCTMEEVDMISLQ	KLQGSV
HC3	SHAF	PIYIKTRVNVTHKEEIIITPIEVAIEDMQKKTQELAFATHQDPADPKMLQ	MLQ	VLQGSV
HC5	MHAF	PIYIKTRISVIQKEEFVITPIEVAIEDMKKKTQLAVAINQEPDAKMLQ	MLQ	VLQGSV

## Coiled-Coil 2

HC2A	SVQVNAGPLAYARAF	LDDTNTKRYPDNKVKLLKEVFRQFVEACGQAI	AVNERLIKEDQLE
KIAA	SVQVNAGPLAYARAF	LDDTNTKRYPDNKVKLLKEVFRQFVEACGQAI	AVNERLIKEDQLE
rat	SVQVNAGPLAYARAF	LDDTNTKRYPDNKVKLLKEVFRQFVEACGQAI	AVNERLIKEDQLE
HC4	SVQVNAGPLAYARAF	LND SQASKYPKKVSELKDMFRKFIQACSI	AI ELNERLIKEDQVE
HC1	SVKVNAGPMAYARAF	LEETNAKKYPDNQVKLLKEIFRQFADACGQAI	AVNERLIKEDQLE
HC3	GTTVNQGP	LEVAQVFLSEIPSDPKLFRHHNKLRLCFKDFTKRCEDAI	RKNKSLIGPVQKE
HC5	GATVNQGP	LEVAQVFLAEIPADPKLYRHHNKLRLCFKEFIMRCGEAVE	EKNKRLITADQRE

## Coiled-Coil 2

HC2A	YQEEMKANYREMAKELSEIMHE	QICPLEEKTS-VLPNSLHIFNAISGTPPTSTMVHG	MTSS
KIAA	YQEEMKANYREMAKELSEIMHE	QLG-----	
rat	YQEEMKANYREIRKELSDIIVER	ICPGEDKRA TKFPAHLQRHQRDTNKHSGSRVDQ	FILS
HC4	YHEGLKSNFRDMVKELSDI	IHEQILQEDTMHSPWMSNTLHVFC	AI SG TSSDRGYGSPRYA
HC1	YQEELRSHYK DMLSELSTVMNE	QITGRDDL SK---RGVDQTCTRVISKATPALPTV	SISS
HC3	YQRELG----	KLSS-----PZ-----	
HC5	YQOELKKNYNKLKENLRPMIER	KIPELYKPIFRVESQKRDSFHRSSFRKCETQLS	QGSZ-

## PBM

HC2A	SSVVZ-----
KIAA	-----
rat	CVTLPHEPHVGTCTCFVMCKLRTTFRANHWFCQAQEEAMGNGREKEPWTVI FNSRFYRSWGK
HC4	EVZ-----
HC1	SAEVZ-----
HC3	-----
HC5	-----

HC2A	-----
KIAA	-----
rat	VHIFF
HC4	-----
HC1	-----
HC3	-----
HC5	-----

Fig. 5A (cont.)

	A	B
CLASP-1	YRVAFYGQ	GFEEEEEGKEYIYKEP
KIAA1058	FRVAFFGQ	AAQYQFTDSETDVEGFFEDEDGKEYIYKEP
CLASP-2		FEDEDGKEYIYKEP
CLASP-6	FRVAFFGQ	GFFEDEDGKEYIYKEP
CLASP-4	FRVAFYGQ	SFFEEEDGKEYIYKEP
DOCK180	FAVGYYGQ	GFPTFLRGKVFIYRGKEYERRED
DOCK2	FAVGYYGQ	GFPSFLRNKVFYIYRGKEYERRED
DOCK3	FRVGIFYGR	KFPFFLRNKEYVCRGH
KIAA0716	FRVGIFYGK	KFPFFLRNKEFVCRGH
CLASP-3	FRVGIFYGT	KFGDLDEQEFVYKEP
CONSENSUS	F V FYG	KEY K
	YF	O F R

TRG	PKLTPLSEISQRL	LLKLYSDKFGSENVKMIODSGKVNPKDLDSKFAY	YIQVTHVTPFFDEKE
CLASP-1	PKLTGLSEISQRL	LLKLYADKFGADNVKIIQDSNKNVNPKDLDPKYAY	YIQVTYVTPFFEEKE
CLASP-2	PKLTPLSEISQRL	LLKLYSDKFGSENVKMTQDSGKVNPKDLDSKYAY	YIQVTHVIPFFDEKE
CLASP-4	PKLTGLSEISLRL	VKLYGEKFGTENVKIIQSDKVNAKELDPKYAH	YIQVTYVVKPYFDQKE
CLASP-3	PAITKLAEIFSHR	LEGFYGERFGEDVVEVIKDSNPVDKCKLDPNKAY	YIQITYVEPYFDTYE
KIAA0716	HDYERLEAFQQR	MLNEFPFAIA-----MQHANQPDETIFQAEAQ	YLQIYAVTPIPEBQE
DOCK3	HDYERLEAFQQR	MLSEFPQAVA-----MQHPNHPDDAILQCDAQ	YLQIYAVTPIPDYVD
DOCK2		FQMQLMTQFPNAEK-----MNTTSAPGDDVKNAPGQ	YIQCFTVQPVLDDEHP
DOCK180	EYERREDFQMQL	MTQFPNAEK-----MNTTSAPGDDVKNAPGQ	YIQCFTVQPVLDDEHP
CONSENSUS	L	L Y	YIQ+ V P D
		M F	L E

	D				E			
CLASP-1	RTIL	TTSHL	FPYV	KKRIQVISQSSTELN	PIEVAIDEM	SRKVSELN		
TRG	RTIL	TAIHCF	FPYV	KKRIPVMYQHHTDLN	PIEVAIDEM	SKKVAELH		
KIAA1058	RTIL	TAIHCF	FPYV	KKRIPVMYQHHTDLN	PIEVAIDEM	SKKVAELR		
CLASP-2	RTIL	TAIHCF	FPYV	KKRIPVMYQHHTDLN	PIEVAIDEM	SKKVAELR		
CLASP-6	RTIL	TAIHCF	FPYV	KKRIPFMYQHHTDLN	PIEV: HDEM	SKKVAELR		
CLASP-4	RTIL	TTSNS	FPYV	KKRIPINCEQQINLKP	IDVATDEI	KDKTAELO		
CLASP-3	KTIL	TTSHAF	FPYIK	TRVNVTHKEEII	LTPIEVAIEDM	QKKTQELA		
CLASP-5	NTVL	TTMHAF	FPYIK	TRISVIQKEEFVLT	PIEVAIEDM	KKKTLQLA		
KIAA0716	RTSL	YLVQSL	PGISR	WFEVEKREVVEMS	PLENAIEV	LENKNQOLK		
DOCK2	RTSF	VTAAYKL	PGIL	RWFEVVHMSQTTIS	PLENAIETM	STANEKIL		
DOCK3	RTTL	TLTHSL	PGISR	WFEVERRELVEVS	PLENAIQVV	ENKNQELR		
DOCK180	RTSF	VTAAYKL	PGIL	RWFEVVHMSQTTIS	PLENAIETM	STANEKIL		
CONSENSUS	RT L		FP V	+ V	+ P+E	AI+ M	+L	
	F		L L			+	I	

[illegible]

DOCK2=KIAA0209  
DOCK3=KIAA0299  
CLASP2variant=KIAA1058

FIG. 5B (cont.)

1  
A

2 32  
GTT TTA CAC CAT CAC CAA AAC CCA GAA TTT TAT GAT GAG ATT AAA ATA GAG TTG CCC ACT  
val leu his his his gln asn pro glu phe tyr asp glu ile lys ile glu leu pro thr

62 92  
CAG CTG CAT GAA AAG CAC CAC CTG TTG CTC ACA TTC TTC CAT GTC AGC TGT GAC AAC TCA  
gln leu his glu lys his his leu leu leu thr phe phe his val ser cys asp asn ser

122 152  
AGT AAA GGA AGC ACG AAG AAG AGG GAT GTC GTT GAA ACC CAA GTT GGC TAC TCC TGG CTT  
ser lys gly ser thr lys lys arg asp val val glu thr gln val gly tyr ser trp leu

182 212  
CCC CTC CTG AAA GAC GGA AGG GTG GTG ACA AGC GAG CAG CAC ATC CCG GTC TCG GCG AAC  
pro leu leu lys asp gly arg val val thr ser glu gln his ile pro val ser ala asn

242 272  
CTT CCT TCG GGC TAT CTT GGC TAC CAA GAG CTT GGG ATG GGC AGG CAT TAT GGT CCG GAA  
leu pro ser gly tyr leu gly tyr gln glu leu gly met gly arg his tyr gly pro glu

302 332  
ATT AAA TGG GTA GAT GGA GGC AAG CCA CTG CTG AAA ATT TCC ACT CAT CTG GTT TCT ACA  
ile lys trp val asp gly gly lys pro leu leu lys ile ser thr his leu val ser thr  
ref 1.1, 1.2 and 1.3

362 392  
GTG TAT ACT CAG GAT CAG CAT TTA CAT AAT TTT TTC CAG TAC TGT CAG AAA ACC GAA TCT  
val tyr thr gln asp gln his leu his asn phe phe gln tyr cys gln lys thr glu ser

422 452  
GGA GCC CAA GCC TTA GGA AAC GAA CTT GTA AAG TAC CTT AAG AGT CTG CAT GCG ATG GAA  
gly ala gln ala leu gly asn glu leu val lys tyr leu lys ser leu his ala met glu

482 512  
GGC CAC GTG ATG ATC GCC TTC TTG CCC ACT ATC CTA AAC CAG CTG TTC CGA GTC CTC ACC  
gly his val met ile ala phe leu pro thr ile leu asn gln leu phe arg val leu thr

542 572  
AGA GCC ACA CAG GAA GAA GTC GCG GTT AAC GTG ACT CGG GTC ATT ATT CAT GTG GTT GCC  
arg ala thr gln glu glu val ala val asn val thr arg val ile ile his val val ala

602 632  
CAG TGC CAT GAG GAA GGA TTG GAG AGC CAC TTG AGG TCA TAT GTT AAG TAC GCG TAT AAG  
gln cys his glu glu gly leu glu ser his leu arg ser tyr val lys tyr ala tyr lys

662 692  
GCT GAG CCA TAT GTT GCC TCT GAA TAC AAG ACA GTG CAT GAA GAA CTG ACC AAA TCC ATG  
ala glu pro tyr val ala ser glu tyr lys thr val his glu glu leu thr lys ser met

FIG. 4A



722 752  
ACC ACG ATT CTC AAG CCT TCT GCC GAT TTC CTC ACC AGC AAC AAA CTA CTG AGG TAC TCA  
thr thr ile leu lys pro ser ala asp phe leu thr ser asn lys leu leu arg tyr ser

782 812  
TGG TTT TTC TTT GAT GTA CTG ATC AAA TCT ATG GCT CAG CAT TTG ATA GAG AAC TCC AAA  
trp phe phe phe asp val leu ile lys ser met ala gln his leu ile glu asn ser lys

842 |Cadherin Cleavage| 872  
GTT AAG TTG CTG CGA AAC CAG AGA TTT CCT GCA TCC TAT CAT CAT GCA GCG GAA ACC GTT  
val lys leu leu arg asn gln arg phe pro ala ser tyr his his ala ala glu thr val

902 932  
GTA AAT ATG CTG ATG CCA CAC ATC ACT CAG AAG TTT GGA GAT AAT CCA GAG GCA TCT AAG  
val asn met leu met pro his ile thr gln lys phe gly asp asn pro glu ala ser lys

962 992  
AAC GCG AAT CAT AGC CTT GCT GTC TTC ATC AAG AGA TGT TTC ACC TTC ATG GAC AGG GGC  
asn ala asn his ser leu ala val phe ile lys arg cys phe thr phe met asp arg gly  
ref 2.1 ↓

1022 1052  
TTT GTC TTC AAG CAG ATC AAC AAC TAC ATT AGC TGT TTT GCT CCT GGA GAC CCA AAG ACC  
phe val phe lys gln ile asn asn tyr ile ser cys phe ala pro gly asp pro lys thr

1082 1112  
CTC TTT GAA TAC AAG TTT GAA TTT CTC CGT GTA GTG TGC AAC CAT GAA CAT TAT ATT CCG  
leu phe glu tyr lys phe glu phe leu arg val val cys asn his glu his tyr ile pro

1142 1172  
TTG AAC TTA CCA ATG CCA TTT GGA AAA GGC AGG ATT CAA AGA TAC CAA GAC CTC CAG CTT  
leu asn leu pro met pro phe gly lys gly arg ile gln arg tyr gln asp leu gln leu

1202 1232  
GAC TAC TCA TTA ACA GAT GAG TTC TGC AGA AAC CAC TTC TTG GTG GGA CTG TTA CTG AGG  
asp tyr ser leu thr asp glu phe cys arg asn his phe leu val gly leu leu leu arg

1262 1292  
GAG GTG GGG ACA GCC CTC CAG GAG TTC CGG GAG GTC CGT CTG ATC GCC ATC AGT GTG CTC  
glu val gly thr ala leu gln glu phe arg glu val arg leu ile ala ile ser val leu  
ref 3.1 ↓

1322 1352  
AAG AAC CTG CTG ATA AAG CAT TCT TTT GAT GAC AGA TAT GCT TCA AGG AGC CAT CAG GCA  
lys asn leu leu ile lys his ser phe asp asp arg tyr ala ser arg ser his gln ala

1382 1412/471  
AGG ATA GCC ACC CTC TAC CTG CCT CTG TTT GGT CTG CTG ATT GAA AAC GTC CAG CGG ATC  
arg ile ala thr leu tyr leu pro leu phe gly leu leu ile glu asn val gln arg ile

1442 1472  
AAT GTG AGG GAT GTG TCA CCC TTC CCT GTG AAC GCG GGC ATG ACC GTG AAG GAT GAA TCC  
asn val arg asp val ser pro phe pro val asn ala gly met thr val lys asp glu ser

1502 1532

Fig. 6A (cont.)

CTG GCT CTA CCA GCT GTG AAT CCG CTG GTG ACG CCG CAG AAG GGA AGC ACC CTG GAC AAC  
leu ala leu pro ala val asn pro leu val thr pro gln lys gly ser thr leu asp asn  
ref 4.1 and 4.2

1562 1592  
AGC CTG CAC AAG GAC CTG CTG GGC GCC ATC TCC GGC ATT GCT TCT CCA TAT ACA ACC TCA  
ser leu his lys asp leu leu gly ala ile ser gly ile ala ser pro tyr thr thr ser

1622 1652  
ACT CCA AAC ATC AAC AGT GTG AGA AAT GCT GAT TCG AGA GGA TCT CTC ATA AGC ACA GAT  
thr pro asn ile asn ser val arg asn ala asp ser arg gly ser leu ile ser thr asp  
ref 5.1 and 5.2

1682 1712  
TCG GGT AAC AGC CTT CCA GAA AGG AAT AGT GAG AAG AGC AAT TCC CTG GAT AAG CAC CAA  
ser gly asn ser leu pro glu arg asn ser glu lys ser asn ser leu asp lys his gln

1742 1772  
CAA AGT AGC ACA TTG GGA AAT TCC GTG GTT CGC TGT GAT AAA CTT GAC CAG TCT GAG ATT  
gln ser ser thr leu gly asn ser val val arg cys asp lys leu asp gln ser glu ile

1802 1832  
AAG AGC CTA CTG ATG TGT TTC CTC TAC ATC TTA AAG AGC ATG TCT GAT GAT GCT TTG TTT  
lys ser leu leu met cys phe leu tyr ile leu lys ser met ser asp asp ala leu phe

1862 1892  
ACA TAT TGG AAC AAG GCT TCA ACA TCT GAA CTT ATG GAT TTT TTT ACA ATA TCT GAA GTC  
thr tyr trp asn lys ala ser thr ser glu leu met asp phe phe thr ile ser glu val  
ref 6.1

1922 1952  
TGC CTG CAC CAG TTC CAG TAC ATG GGG AAG CGA TAC ATA GCC AGG AAC CAG GAG GGG TTG  
cys leu his gln phe gln tyr met gly lys arg tyr ile ala arg asn gln glu gly leu

1982 2012  
GGA CCC ATA GTT CAT GAT CGA AAG TCT CAG ACA TTG CCT GTT TCC CGT AAC AGA ACA GGA  
gly pro ile val his asp arg lys ser gln thr leu pro val ser arg asn arg thr gly

2042 2072  
ATG ATG CAT GCC AGA TTG CAG CAG CTG GGC AGC CTG GAT AAC TCT CTC ACT TTT AAC CAC  
met met his ala arg leu gln gln leu gly ser leu asp asn ser leu thr phe asn his

2102 2132  
AGC TAT GGC CAC TCG GAC GCA GAT GTT CTG CAC CAG TCA TTA CTT GAA GCC AAC ATT GCT  
ser tyr gly his ser asp ala asp val leu his gln ser leu leu glu ala asn ile ala

2162 2192  
ACT GAG GTT TGC CTG ACA GCT CTG GAC ACG CTT TCT CTA TTT ACA TTG GCG TTT AAG AAC  
thr glu val cys leu thr ala leu asp thr leu ser leu phe thr leu ala phe lys asn  
ref 7.1

2222 2252  
CAG CTC CTG GCC GAC CAT GGA CAT AAT CCT CTC ATG AAA AAA GTT TTT GAT GTC TAC CTG  
gln leu leu ala asp his gly his asn pro leu met lys lys val phe asp val tyr leu

2282 2312

FIG. 6A (cont.)

TGT TTT CTT CAA AAA CAT CAG TCT GAA ACG GCT TTA AAA AAT GTC TTC ACT GCC TTA AGG  
cys phe leu gln lys his gln ser glu thr ala leu lys asn val phe thr ala leu arg

2342

2372

TCC TTA ATT TAT AAG TTT CCC TCA ACA TTC TAT GAA GGG AGA GCG GAC ATG TGT GCG GCT  
ser leu ile tyr lys phe pro ser thr phe tyr glu gly arg ala asp met cys ala ala

2402

2432

CTG TGT TAC GAG ATT CTC AAG TGC TGT AAC TCC AAG CTG AGC TCC ATC AGG ACG GAG GCC  
leu cys tyr glu ile leu lys cys cys asn ser lys leu ser ser ile arg thr glu ala

2462

2492

TCC CAG CTG CTC TAC TTC CTG ATG AGG AAC AAC TTT GAT TAC ACT GGA AAG AAG TCC TTT  
ser gln leu leu tyr phe leu met arg asn asn phe asp tyr thr gly lys lys ser phe

2522

2552

GTC CGG ACA CAT TTG CAA GTC ATC ATA TCT GTC AGC CAG CTG ATA GCA GAC GTT GTT GGC  
val arg thr his leu gln val ile ile ser val ser gln leu ile ala asp val val gly

2582

2612

ATT GGG GAA ACC AGA TTC CAG CAG TCC CTG TCC ATC ATC AAC AAC TGT GCC AAC AGT GAC  
ile gly glu thr arg phe gln gln ser leu ser ile ile asn asn cys ala asn ser asp

2642

2672

GGG CTT ATT AAG CAC ACC AGC TTC TCC TCT GAT GTG AAG GAC TTA ACC AAA AGG ATA CGC  
arg leu ile lys his thr ser phe ser ser asp val lys asp leu thr lys arg ile arg

2702

2732

ACG GTG CTA ATG GCC ACC GCC CAG ATG AAG GAG CAT GAG AAC GAC CCA GAG ATG CTG GTG  
thr val leu met ala thr ala gln met lys glu his glu asn asp pro glu met leu val

2762

2792

GAC CTC CAG TAC AGC CTG GCC AAA TCC TAT GCC AGC ACG CCC GAG CTC AGG AAG ACG TGG  
asp leu gln tyr ser leu ala lys ser tyr ala ser thr pro glu leu arg lys thr trp

2822

2852

1xxxxxxxxxxxxxxxx Predicted

CTC GAC AGC ATG GCC AGG ATC CAT GTC AAA AAT GGC GAT CTC TCA GAG GCA GCA ATG TGC  
leu asp ser met ala arg ile his val lys asn gly asp leu ser glu ala ala met cys

Transmembrane Domain xxxxxxxxxxxxxxxxxxxxxxxxxxxxxxxx|

TAT GTC CAC GTA ACA GCC CTA GTG GCA GAA TAT CTC ACA CGG AAA GGC GTG TTT AGA CAA  
tyr val his val thr ala leu val ala glu tyr leu thr arg lys gly val phe arg gln

2942

2972

GGA TGC ACC GCC TTC AGG GTC ATT ACC CCA AAC ATC GAC GAG GAG GCC TCC ATG ATG GAA  
gly cys thr ala phe arg val ile thr pro asn ile asp glu glu ala ser met met glu  
ref 8.1 ↓

3002

3032

GAC GTG GGG ATG CAG GAT GTC CAT TTC AAC GAG GAT GTG CTG ATG GAG CTC CTT GAG CAG  
asp val gly met gln asp val his phe asn glu asp val leu met glu leu leu glu gln

3062

3092

TGC GCA GAT GGA CTC TGG AAA GCC GAG CGC TAC GAG CTC ATC GCC GAC ATC TAC AAA CTT  
cys ala asp gly leu trp lys ala glu arg tyr glu leu ile ala asp ile tyr lys leu

Fig. 6A (cont.)

ref 9.1

3122 ATC ATC CCC ATT TAT GAG AAG CGG AGG GAT TTC TTT GAA GAT GAA GAT GGA AAG GAG TAT  
ile ile pro ile tyr glu lys arg arg asp phe phe glu asp glu asp gly lys glu tyr

3152

3182 ATT TAC AAG GAA CCC AAA CTC ACA CCG CTG TCG GAA ATT TCT CAG AGA CTC CTT AAA CTG  
ile tyr lys glu pro lys leu thr pro leu ser glu ile ser gln arg leu leu lys leu

3212

ref 10.1

3242 TAC TCG GAT AAA TTT GGT TCT GAA AAT GTC AAA ATG ATA CAG GAT TCT GGC AAG GTC AAC  
tyr ser asp lys phe gly ser glu asn val lys met ile gln asp ser gly lys val asn

3272

3302 CCT AAG GAT CTG GAT TCT AAG TAT GCA TAC ATC CAG GTG ACT CAC GTC ATC CCC TTC TTT  
pro lys asp leu asp ser lys tyr ala tyr ile gln val thr his val ile pro phe phe

3332

3362 GAC GAA AAA GAG TTG CAA GAA AGG AAA ACA GAG TTT GAG AGA TCC CAC AAC ATC CGC CGC  
asp glu lys glu leu gln glu arg lys thr glu phe glu arg ser his asn ile arg arg

3392

3422 TTC ATG TTT GAG ATG CCA TTT ACG CAG ACC GGG AAG AGG CAG GGC GGG GTG GAA GAG CAG  
phe met phe glu met pro phe thr gln thr gly lys arg gln gly gly val glu glu gln

3452

ref 11.1

3482 TGC AAA CGG CGC ACC ATC CTG ACA GCC ATA CAC TGC TTC CCT TAT GTG AAG AAG CGC ATC  
cys lys arg arg thr ile leu thr ala ile his cys phe pro tyr val lys lys arg ile

3512

3542 CCT GTC ATG TAC CAG CAC CAC ACT GAC CTG AAC CCC ATC GAG GTG GCC ATT GAC GAG ATG  
pro val met tyr gln his his thr asp leu asn pro ile glu val ala ile asp glu met

3572

1xxxxxxx Coiled-coil 1 xxxxxx

3602 xxxxxxxx Coiled coil 1 cont'd xxxx 3632 xxxxxxxxxxxxxxxxxxxxxxxxxxxxxxxxxxxx  
AGT AAG AAG GTG GCG GAG CTC CGG CAG CTG TGC TCC TCG GCC GAG GTG GAC ATG ATC AAA  
ser lys lys val ala glu leu arg gln leu cys ser ser ala glu val asp met ile lys

3662 xxxxxxxxxxxxxxxxxxxxxxxxx 1 3692 GTT CAG GTC AAT GCT GGC CCA CTA GCA TAT  
CTG CAG CTC AAA CTC CAG GGC AGC GTG AGT val gln val asn ala gly pro leu ala tyr

3722

3752 GCG CGA GCT TTC TTA GAT GAT ACA AAC ACA AAG CGA TAT CCT GAC AAT AAA GTG AAG CTG  
ala arg ala phe leu asp asp thr asn thr lys arg tyr pro asp asn lys val lys leu

3782

3812 1xxxxxxxxxxxxxxxxxxxxxx  
CTT AAG GAA GTT TTC AGG CAA TTT GTG GAA GCT TGC GGT CAA GCC TTA GCG GTA AAC GAA  
leu lys glu val phe arg gln phe val glu ala cys gly gln ala leu ala val asn glu

3842 xxxxxxxx Coiled coil 2 xxxxxxxxxx 3872 xxxxxxxxxxxxxxxxxxxxxxxxxxxxxxxxxxxx  
CGT CTG ATT AAA GAA GAC CAG CTC GAG TAT CAG GAA GAA ATG AAA GCC AAC TAC AGG GAA  
arg leu ile lys glu asp gln leu glu tyr gln glu glu met lys ala asn tyr arg glu

3902 xxxxxxxxxxxxxxxxxxxxxxxxxxxxxxxxxxxx 3932 xxxl

FIG. 6A(cont.)

ATG GCG AAG GAG CTT TCT GAA ATC ATG CAT GAG CAG ATC TGC CCC CTG GAG GAG AAG ACG  
met ala lys glu leu ser glu ile met his glu gln ile cys pro leu glu glu lys thr

3962

3992

AGC GTC TTA CCG AAT TCC CTT CAC ATC TTC AAC GCC ATC AGT GGG ACT CCA ACA AGC ACA  
ser val leu pro asn ser leu his ile phe asn ala ile ser gly thr pro thr ser thr

4022

lxxxx PBM xxxxxl

ATG GTT CAC GGG ATG ACC AGC TCG TCT TCG GTC GTG TGA TTA CAT CTC ATG GCC CGT GTG  
met val his gly met thr ser ser ser ser val val STP

4082

4112

TGG GGA CTT GCT TTG TCA TTT GCA AAC TCA GGA TGC TTT CCA AAG CCA ATC ACT GGG GAG

4142

4172

ACC GAG CAC AGG GAG GAC CAA GGG GAA GGG GAG AGA AAG GAA ATA AAG AAC AAC GTT ATT

4202

4232

TCT TAA CAG ACT TTC TAT AGG AGT TGT AAG AAG GTG CAC ATA TTT TTT TAA ATC TCA CTG

4262

4292

GCA ATA TTC AAA GTT TTC ATT GTG TCT TAA CAA AGG TGT GGT AGA CAC TCT TGA GCT GGA

4322

4352

CTT AGA TTT TAT TCT TCC TTG CAG AGT AGT GTT AGA ATA GAT GGC CTA CAG AAA AAA AAG

4382

4412

GTT CTG GGA TCT ACA TGG CAG GGA GGG CTG CAC TGA CAT TGA TGC CTG GGG GAC CTT TTG

ref 13.1

4442

4472

CCT CGA CTC GTG CCG GAA ATC TGA TCG TAA TCA GGG TAC AGA ACT TAC TAG TTT TGT CTA

4502

4532

GGA GTA TGT TGT ATG ACT AGG ATT TGT GCT ATT ATC TCA TTC AAC AAC ATA GAG CAA GAA

4562

4592

TAG TGA GCT AAC TGA GCT AGA CAC TCA ATT AAT CCG CTA CTG GCT TCA AGT CAG AAC TTT

4622

4652

GTC ATT AAT CAT CGA CTC CGG GAC GGT CAT ATA TGT ATT ACA TTT CTA CAT TTT TAA TAC

4682

4712

TCA CAT GGG CTT ATG CAT TAA GTT TAA TTG TGA TAA ATT TGT GCT GGT CCA GTA TAT GCA

4742

4772

ATA CAC TTT AAT GGT TTA TTC TTG TCA TAA AAA TGT GCA ATA TGG AGA TGT ATA CAA GTC

4802

TTT ACT

Flt. Lt. (cont.)

## BAC sequences of Human CLASP 2

### Ref 1.1

Sequence of BAC4 using primer HC2AS2, which spans nucleotides 327-346 of the cDNA. Exon sequence is underlined and represents nucleotides 356-375.

TTTCTACAGNGTNTACTCAGGTATGTGCTCCTTCAACAAAATTAGCAGTTGCTGCTCTG  
TGACAAAGTTTGCACCATTTTGTCAAGAAGAAAAAATCCTAATGTGTTATATTACTATA  
TTTTTACTCTATAGATCTTTTTTCTAAAGAAAGAAAGTACAACCTGAAGTGCTTATATGTA  
TTCATATAAATGACTAGTACAAGCATCATTTTGTCAACAGATTTCCCCTTTCATTGGAGG  
ATCTTCTTGATGTTATTTGTACACGATCAATTTTGTAGTCTTAATAAGATGAGGCTGGGTG  
TGGTGGCTCACACCTGTAATCCTAGCATTTTGGAGGCCAAGGTGGGCAGATCACTTTAG  
CCCAGGGGTTTGAGACCAGCCTGGCCAACATGGCAAAACCTTGTCTCTACAAAAATAC  
NAAAATTATCCAGGCATGGTGATGTGTGCCTGTAGTCCCAACTNCCTAGGAGGCTAGG  
GGTAGGGGGGATTTGCAAGAGGCTGGGAGGGGTCAAAGCCCNAANTGAGCCATTGGTNC  
ATGTCACTTGGACCCCAAGCNGGGGGNGANCAAGAGCAAAGGACTNNTGTNNTTTAN  
AAAAAAAACCGGGCTACCATACNNACCAACCCNCNNACCTACCCNACCTTTCCANNTT  
AAAANAAGGCTTTGNCTTGCANAGGAAAANCAAAATNNCC

### Ref 1.2

Sequence of BAC26 using primer HC2AS2, which spans nucleotides 327-346 of the cDNA. Exon sequence is underlined and represents nucleotides 351-375.

TCTGGTTTCTACAGTGTATACTNAGGTATGTGCTCCTTNAACAAAATTAGCAGTTGCTG  
CTCTGTGACAAAGTTTGCACCATTTTGTCAAGAAGAAAAAATCCTAATGTGTTATATTA  
CTATATTTTTTACTCTATAGATCTTTTTTCTAAAGAAAGAAAGTACAACCTGAAGTGCTTAT  
ATGTATTCATATAAATGACTAGTACAAGCATCATTTTGTCAACAGATTTCCCCTTTCATT  
GGAGGATCTTCTTGATGTTATTTGTACACGATCAATTTTGTAGTCTTAATAAGATGAGGC  
TGGGTGTGGTGGCTCACACCTGTAATCCTAGCATTTTGGAGGCCAAGGTGGGCAGATC  
ACTTTAGCCCAGGGGTTTGAGACCAGCCTGGCCAACATGGCAAAACCTTGTCTCTACA  
AAAATACAAAAATTATCCAGGCATGGTGATGTGTGCCTGTAGTCCCAGCTACCTAGGA  
GGCTAGGGTAGGGGGGATTTGCAAGAGGCTNGGAGGTCAAGGCCCGCAGTGAGCCATGG  
TCATGTCACTGCACCCCCAGCCAGGGCCGACAGGAGCAAGACTNTTGTNTCAAAAAAA  
AACAGNAACCAACANCCAACAACAACNACCTTTCNGCAAAANAAGCTTGCTNCA  
ANGAAACCAAAATGNCTTCTTNTTTTCCCCCN

### Ref 1.3

Sequence of BAC26 using primer HC2AS2, which spans nucleotides 327-346 of the cDNA. Exon sequence is not found within this sequence. This sequence most likely represent intron sequence since this sequence matches the intron sequence found in the previous two BAC sequences.

AGNNNNNCCCNCTACNCCACTTTTAACCTTTTGAAAACACAGTGTTTNTCTCAANTATGC  
GCTCCTTCACATATTAGCAGTTGCTGCTCTGTGACATAGTTGCACCATNTGTCAAGAAG  
AAAAAATCCTAAGTGTNATATCACTATATNNNTACTCTATAGATCTTNTCTAAAGAAAG  
AAAGTCAACTGATGTGCTTATATGTATNCATATAAATGACTAGTACATGCATCATTTTG  
CAACAGATNTCTCCTCACATTGGAGGATCTTCTNGANGNATTCGACACGATNANTATTA  
GTCTNAATAAGATGANGCTGGTGTGGNGGTACACTGNATCTAGCATNTGGANGCATGT

Fig. 6A (cont.)



GGCAGACACTTANCCNCGGTNGAGACAGCTGTCAGTGNCAACTGTCTCTNTAAANCA  
 AANNCTCCGCNNGGNGATGGGCTGAGCCAGTCCTAGNNGCTAGNTAGNGATGNNGAGN  
 TGTNGCACGNCGAGNGAGCATGNTCTGTACTGACTCATCAGGCGNCNACACGNTCTGT  
 TCNAAAACATAACCACACACACTCNCACCTNCGCAAAATTGCTCTNNAANATGCTTNT  
 TTCACACNGNTNCAATCNCTATATNNTCTTCTATTCTNACGTNTNATTANNATCTTN  
 CNCTGCANAAACNATNCGNCCACCTNNANNACCTTANGCTTNGTTTCACGCTTATAGCTC  
 CCCTACACNTNNCAGCNNTTNCNNGTGAAGGGCCNCCCGAATCTACGANCACTCTC  
 TCCGTATATNGCCTCGGTCCANCGCCATCTGCTGTNTNCTCNCNCTNCGCNNTTNANCNG  
 TNCGCTATCTCTNNNCCGGATCCNCCCATATNNTNNCTCTACTTANAGCGTAANNTNT  
 NCNCACTANTCACAACCTTNTNCTNNAACTCTATCTNCTCCTCTCTACCACCTCACT  
 TACTACCTNTTCACNCANTCTCCTTCNCTNTCCACTGATCTCCACATAGCTGCTNTACTC  
 GCCANTTTATCATATNCACACNCTCTACGCTNNNTNT

#### Ref 2.1

Sequence of BAC4 using primer HC2S1, which spans nucleotides 1107-1126 of the cDNA. Exon sequence is underlined and represents nucleotides 1079-1097.

CTTGTATTNAAAGAGGGTCTGCAGGAAGAAGTGTGTAGTCATAAATACCTCACTGGAT  
 ATTTTATACAGGATTCTAAAAAACCTATTAGCAATAGTATGCTAGAAATAGTCATTAGC  
 TTCTTGACCTTCTTAGAACTGCACACTCTATTGCACTGTACAGATTTTCAGGATGGCTGC  
 AGGGATTGATTTGAAAACCTAAGGACACATTTCAATAAACAATGTCTTCAATTGATTTTT  
 AGGGCTCCTCCTACTTCAATGAAGGACTTCAGGTAGCTTATAATTACAGACACAGGCTC  
 AATACAATAAAAAAATTAGTAAGGCAGAGCTTTAAAAAAGGAAAAAGATAA  
 TTCTACCAGAGAAAGGCTACATGGTGACTTCTGTTACCAGTAACAACCCCGCACTACC  
 TTTGGGTCTCCAGGAGCAAAACAGCTAATGTAGTTGTTGATCTGCTTGAAGACAAAGC  
 CCCTGTCCATGAAGGTGAAACATCTCTGTGGAGGAAACAAGCAAAAAAGTTATTTCA  
 GGTCCAAACATTTTCGGAAATTTGGATTCAAAGCAGGCATTTATTGCTAATAAGTTTATC  
 CACTGACATAAAAAACATGCCTTCAACATTGCCAGAGCACCTACTCTATTNTAGTCNCN

#### Ref 3.1

Sequence of BAC4 using primer C96AS, which spans nucleotides 1443-1452 of the cDNA. Exon sequence is underlined and represents nucleotides 1370-1422.

AATCAGCAGACCAAACAGAGGCAGGTAGAGGGTGGCTATCCTTGCCTGATGGCTCTGA  
 AAAGAAGACACACATGGTAAGTTTGACCCAGGATTCTGAGAACCGAACTAAGTTGGTG  
 CTGACCATCTCCTTTATTTGGATCCTTCCCTATAAAGACAGATATTTGATTTTAGTCCCAA  
 AATAGAGCAAAATCTTAGTGCTGTTACCATGAATTTTCTAACTGATTACTTTCTTTACAC  
 CACTTAAAATAAAGGACATTATCAATGCACATTCCTTCCATTGGGGACCACTCACCTT  
 GAAGCATATCTGTCATCAAAAGAATGCTTTATCAGCAGGTTCTTGAGCACACTGATGGC  
 GATCAGACGGACCTCCCGGAACTCCTGGAGGGCTGTCCCCACCTCCCTNAGTAACAGT  
 CCCACCAAGAAGTGGTTTCTGCAGAACTCATCTGTTAATGAGTAGTCAAGCTGGGAGG  
 TCTGAAATGAGGATAGAACTACTTTGNGTTAGGAAAGATGCAATGCTCTTTTGAATA  
 AAACAAACAAACCAAACNAACAAAAAATAAGACCCATCCTTNTGNATTTCAA  
 GCCCACCCTGGGGTNGGTCAAAGAGATGATCAGNANTTTGGCNTTNAAATGAAGAAAG  
 AAATNAATTNTCCAGGGGNTGTTCTNCTTTTATGCACANGGAGGGATNTTAANTGAAA  
 ACCAATTTAAATCCAATTNAGGNG

#### Ref 4.1

FIG. 6A (cont.)

Sequence of BAC4 using primer C2AS5, which spans nucleotides 1716-1735 of the cDNA. Exon sequence is underlined and represents nucleotides 1602-1703.

TTCCTTTCTGCAAGGCTGTTCCCGAATCTGTGCTTATGAGAGATCCTCTCGAATCAGCA  
TTTCTCACACTGTTGATGTTTGGAGTTGAGGTTGTATATGGAGAAGCTAAATGGAAATC  
AAGCCAACAATAAAGTTTTATTAAGACAGAACAAAATAAAGATGAGTACTGAACTTTA  
AGGGAAATTGCTTTTATTGCACTTATTTTTTCTGTTAGGAAGTTGGCTCAAGAGTTGCAT  
TCCATTACTTCACCTTTAAAGAACCAGGTCATATACAATGAGATAAAAAGAACTAGT  
CTGAAACATTCAGATGTAAACATCAATTCACCTTGTTAGAAACCACCTTTGATCGCTAAA  
GACTAAATGCATACCTGTTTCAGAATGTGATAGAATGAAGACTTAAAAAAATTAAAAG  
ATAAATCCACCTACAACCTATCAAATCACAAAATTAAACCACACAACAACTTGTAGCA  
TTCAAACCTGGTAATAAACACTGAGGAGCCTACCCAACCTCTGAGGGGGTGTGATGGGGTA  
TTTTAAATTTTCGAGGAGAACACAGTGATATGTGACCTCAGCCAGAAGCTGCTGTTTNA  
GCAGCAGGTTGGTGCTATGCTCCTTTTTGAAGACATATTTGTGAAGCTGGGTATTTTGG  
GGGCCTGCTTATGATAAAAANGGCAAGGTNTTCAATGNAGGGGN

#### Ref 4.2

Sequence of BAC26 using primer C2AS5, which spans nucleotides 1716-1735 of the cDNA. Exon sequence is underlined and represents nucleotides 1602-1703.

TTCCTTTCTGGAAGGCTGTTACCCGAATCTGTGCTTATGAGAGATCCTCTCGAATCAGC  
ATTTCTCACACTGTTGATGTTTGGAGTTGAGGTTGTATATGGAGAAGCTAAATGGAAAT  
CAAGCCAACAATAAAGTTTTATTAAGACAGAACAAAATAAAGATGAGTACTGAACTTT  
AAGGGAAATTGCTTTTATTGCACTTATTTTTTCTGTTAGGAAGTTGGCTCAAGAGTTGC  
ATTCCATTACTTCACCTTTAAAGAACCAGGTCATATACAATGAGATAAAAAGAACTA  
GTCTGAAACATTCAGATGTAAACATCAATTCACCTTGTTAGAAACCACCTTTGATCGCTA  
AAGACTAAATGCATACCTGTTTCAGAATGTGATAGAATGAAGACTTAAAAAAATTAAA  
AGATAAATCCACCTACAACCTATCAAATCACAAAATTAAACCNCACAACAACTTGTAG  
CATTCAAACCTGGTAATAAAACACTGAGGAGCCTACCCAACCTTTGAGGGGGTGTCAATGG  
GGTNTTTTTTAAATTTTTTCGNNGGANANCCAGTGNTATGGTGACCTTCACCCAAGAAGC  
TTGTTTGTNNACCAAGCNAGGTTGNNCTNTGCTCCTTTTTAGAAANACNNTATTTTNNN  
AAATNCTGGNTTTTTTNNGNGGCCCTNCNTTNT

#### Ref 5.1

Sequence of BAC4 using primer C2S6, which spans nucleotides 1686-1705 of the cDNA. Exon sequence is underlined and represents nucleotides 1724-1736.

TTCCTGGATAAGGTAATTGCTTTTACCCAACACAAATGTTTCTTATAATCAATGGATTT  
AGCCCAAAGTAAACGTACTTCATGTTCTAGTGCCTTTTAAGTGTGACCTTTTGTTTTTTT  
CTAAACCACCCGGCTGACCTGGAGTAGGTGATGAGAGCTTTAAGGTTGGGGCCCATTC  
CTTGAAGTGCTCTGATTCCTGTTTCCAGTACCTCAGATCCTGGGCAGGGTTTGCAGTGG  
AGCGTCTTGAGTGAATGGCTCTGGTGGGTTGAACGGGGAGGGACTCAAATGCTGCCC  
ATCTCAATTTCTGTAGTCTTTTTATTATTTATTTATTTTTTGAGACAGAGTCTCGCTCT  
GTCGCCCAGGCTGGAGTACAGCGGCACGATCTCAATTNACTGCAACCTCCGCCTCC:TG  
GGTTCAAACGACTCCTCTGCCTCAGCCTCCCCAGCAGC:TGGGACCACAGGCACAAGCC  
ACCACCGCCCGGCTAATTTTTTGTNTTTTAGTA:GAGAT:GGGGTTTCACCATATTTGGC  
CAGGCTGGGCTCAAACCTCCTGACC:TCGTCAATCCGCNCCCTCGGNCTNCCAAAGTGCTT  
GGGATTNCAGGCNGTGAGCCCACTTACACCTNNGGCAATTCCCTGTNAGTCTTTTTTAC  
CAGAGACACCATCATTCAACACAGCTTTTCCACCCACAA

Fig. 6A (cont.)

### Ref 5.2

Sequence of BAC26 using primer C2S6, which spans nucleotides 1686-1705 of the cDNA. Exon sequence is underlined and represents nucleotides 1712-1736.

TGAGAAGAGCAATTTCTGGATAAGGTAATTGCTTTTACCCAACACAAATGTTTCTTAT  
AATCAATGGATTTAGCCCAAAGTAAACGTACTTCATGTTCTAGTGCCTTTTAAGTGTGA  
CCTTTTGTTTTTTTCTAAACCACCCGGCTGACCTGGAGTAGGTGATGAGAGCTTTAAGG  
TTGGGGCCCATTCCTTGAAGTGCTCTGATTCCTGTTTCCAGTACCTCAGATCCTGGGCA  
GGGTTTGCAGTGGAGCGTCTTGAGTGAATGGCTCTGGTGGGTTGAACGGGGAGGGACT  
CAAAATGCTGCCCATCTCAATTTCTGTAGTCTTTTTATTTATTTATTTATTTTGTGAGAC  
AGAGTCTCGCTCTGTCGCCCAGGCTGGAGTACAGCGGCACGATCTCAATTCAGTGCAA  
CCTCCGNCTCCCTGGGTTCAAACGACTCCTCTGNCTNAGNCTCCC:AGCAGCCTGGGAA  
CCACAGGCTCANGCCACCACGCCCCGGCTAATTNTTGTAATTTTNAAGTAANAAATTGGG  
GGTTCTCACCATNTTGGCCCAAGNCTTGGGCCTAAAAACCTTNCTNACCNTCGNCATTC  
NCNCCCCNACCNTGGGCNCTNCTCAAANGNGCTTGGGGATTANCANNGGCNTTAACC  
CCCCNTATCACCGTGGNCCTTAATTT

### Ref 6.1

Sequence of BAC4 using primer C2S7, which spans nucleotides 1918-1937 of the cDNA. Exon sequence is not found within this sequence. Since the primer is directed against exon sequence we presume that sequence derived from C2S7 is intron sequence.

NAGNGNGGGTTTNAGNCGTTTGAAGCCTGNNACGNGGTGNGTGCTNGAACTCTGTGGG  
CTTTCAGGTAAGTGGGGTATCTGGGAGCCTGCTGTTTGCATTGCTAGTGCATCAGACCAG  
GGCTTTTCTCCTCCCTGTAGCTGCTACTTATACACATAGCTCTAACTGAGATGATTCTCCA  
GACAACTGATGCAGAGCAGCAAAAGCTTCTGCCGTTCTCCCCTTCTAGGAGTGTCTCCT  
TTCTTTGGAAAGAGATCATGAGGGGCTAGATTGTAATGAAGTGAGGCTCAGTGCTTGA  
GCACATCCGGTAAAAGTTCCAATATATTGGTCATAAAGTTTCTCATTCTTTATAGCAGT  
TAATTTCTCTGGCTCATGAGTTTTCTTAGTTTTAATCTGACTTTTAAATTAATGTCTCCA  
GCACCAGTCATATCCCCAGGGCAAACCTCAAAGGCATGAGAGGCCAGACTCGGGTCCTG  
GTCATAGCAACCCCTGTCTAGGGCCTTGGTCCCTGCCTCCGCTTGTGTGCTGTGGCGCA  
GGTCCTATGGGCCCTTAGGAAACAGGACCACCCTGTCGCACCCCCTACAGAGACCAGC  
CAAGTTTGACATTAGATCACCGTAGCAATGTNTGCAAATTCCAGTTTCTTGCTAAAACA  
GGTTAAGCCTTGCAGCCACTTTATCTGTAAGTGGCNGAGGTTTTGACATAAAA

### Ref 7.1

Sequence of BAC4 using primer C2S8, which spans nucleotides 2143-2162 of the cDNA. Exon sequence is underlined and represents nucleotides 2182-2219.

CTCTCGACACGCTGTTTCTATTAACATTGGCGTTTAAGGTTTGTATCAATTTGCTGTTCTG  
NGGTTCTAGTTTTACCTTTCACATTCATTCTGCTTGGTAAGCTCAGTGAGCACAACTTA  
CTATGTTGCATTTTTACTTCAGCAATTATTTTTGTCCCTGTAAGGAAACCATTAATCTTT  
AAATTCCTTTAATGAAATCATTCCACAGTGAATGGCTTGAATGCCCTGAAATAAAATTT  
AACTGGTCAGTGTGTGCTGCGCGCTTGGGTATGGTGGAAACACGGTCTCTGGAGGCAG  
TTAACTCTTGGCTCGAACCTTGAGGATGGTGAATATAGGCACCTAATCAGGCATTTCTG  
CCTTGAATATCTTTAAATATATCCAAATGTTATAGCGTTTAATTAGATTTTTATGTAGAA  
AGGAGCAATAAACACAAGACACATGTTTTTCAGTTTTTTATCTGTTACTGCATTAAATGA

Fig. 6A (cont.)



TAAAAACGTTTTGGAGATAGAAAATGAAAGGGGGTTTTTTTTTTGTCTTGTTTTAAAGTT  
TTAGCAAATAATATTCAAGTAGGTGGAGATGGACTCTTCACCACTCTCCTGTTTTTAGG  
AACCCAATACTTTTTTCATTCTTGCTAAATGATTACTTCCATTTCTAGCATAGAAAAGGA  
GAAAATTGGAATGAGTGTTTATAT

#### Ref 8.1

Sequence of BAC4 using primer C2S9, which spans nucleotides 2992-3011 of the cDNA. Exon sequence is not found within this sequence. Since the primer is directed against exon sequence we presume that sequence derived from C2S9 is intron sequence

CGCTTTNAAATNCCAGCCGCTACTGCGGGGCGNTNAAATTCGAAACGTGTTGTTNTCTGT  
GATGCCTGGCTCTGATTGTGTGGGATTGGTCATCAGTGGCGGTTGGCAGNTGGGGTTCA  
TGGAAGCGGCCATGGGGACTGATGGCAGGCCCTTGGATTGCCACCGCAGAGCCTGGCA  
GTGTCTTTGGTCTGCATTCCTACCGGCGAAGTCTCATTTACCTCACGTGTTATCTCTTG  
GAAAGCATTCCTTTAGCGGGCTGTGTCTACCCTTCCATCCTCTCGTCCAAACTCCCCCTC  
CTTCTCTGTTCTGTCTCCTTCCCATCCTCTTCTCCCCAGTTCTTCTTCCTATGTTCCCTTCT  
CAGTGGTTTCTCTTCCTCTGTTTGACTTTCCAAGGTCATTTTGACTGTTCCCTGCTCCCAA  
CTACAAAGATACTAAAATCTCACCTAACCCTCTTCTTCTTTCTTAATGAAAGAATGTT  
TTCAGTCCATCCCAAATTTGTGTGGACTTCACAAACCTTCTCTAAATGGAGCCTTTTCT  
CTTCCTACTCTTGACTAGNTGGTAAACGCTCCATGTTCTTGGCCAGAACTCCCTGGTGA  
GTAGCGTCACTCCCACTTTCCTGTGCAGAACCAAGCCTCCTAGAAAACCTCTTTGCANC  
TGAGTGGGTTGGGACACGCCCTTTNTTTGGG

#### Ref 9.1

Sequence of BAC4 using primer C2AS10, which spans nucleotides 3276-3295 of the cDNA. Exon sequence is underlined and represents nucleotides 3147-3234.

TTTANACCNATNTATCCGNGTCAGTTANAGGAGTCTCTGAGAAATTTCCGACAGCGGT  
GTGAGTTTGGGTTCCCTTGTAATATACTCCTTTCCATCTTCATCTTCAAAGAATCCCTGT  
GACATAAAGCACAATTAGAGCTATCCCTGAACGTAAGCCCAGGGCTTACCACCTAGGA  
AGCGTTCTTTTATTACAAGGGGGGAAAAAAGGAATGGGTCTAAAAATCCAGCTGAAAT  
GGGCTTTCTGAATGAGAAAGAAAAATGCTAATAACATGAAGTCTAGGTGCAAAGGTAAA  
GGAAAAACACAACATTGCAAACCTTATTCAAGAATGCAGTCATTAAGTGTTGAGTGAAA  
TGAAAGATTTTGGATACAAGACTAAGCTGTCCCAGGGAAGTCTAATGGGAGTCAAGCC  
TGTTTCACTTTCCCAAGAAGCAGAACTCACTANAAAATGATGAGCAGCCCACGACAGG  
CAGGCTCAGAAGTGGACATGCCTCCCTTCTCCTGATGGCTNCCATGCACACAGGATTTT  
ATGGCATGAAGTGAAGCGTTTGGGGGTCTGGAGTAAGTTTAGTAAAAGTTAGGTAAAG  
CTTGTATAAATTGTATTTTGTCTTACCCGATGAGAAAAAAAATATTNAAGACCTGGTA  
GCTTCAATATTCAAGAAAAATATTTTTCATNTCACCCG

#### Ref 10.1

Sequence of BAC4 using primer C2S11, which spans nucleotides 3167-3186 of the cDNA. Exon sequence is underlined and represents nucleotides 3231-3296.

NGNANGTGGAGCCNCGANCCAGGGACAATCTNAACCTNCTTAAACTGTACTCGGATNA  
ATTTGGTTCTGAAAATGTCAAAATGATACAGGATTCTGGCAAGGTATTGACCATGTTTG  
ANAAGTTTCATAGCAATGTAATGTTGTGATNCGATTACATATNATATATTTTAAATG  
TNTATAGAAAAAACACANGAAAAATATTAAGGATTGTTGGCCCGTGAGTGGCAGGTG  
TATNTTCTTNCTGATCCTTTAGNGCTTTCATTACATGCNTGACATTAAAAAAANCTTTA

Fig. 6A (cont.)

TCGCCTAATTTTTGAAACATCTAATTTTACAAAATAATTAACCGTNTGGCCANGNATAT  
TNTCATTTTTAGGNCCAGCTATTTAGAAACTCTGACANAAATGAGGGGCTGTGGCTTNC  
CTNCCTNNACTTGNCCTCTTTTCNNGNATGTACCACATGAACTTGNCNCCTCTTTTCNNC  
TNACCGGGTGGCATGTTANAGGACAGGTTGAAACCNCANTNGGGCNGGANTTNGGTN  
NAATTGGGACACAATGGTACNANGCTCTATNGGAATNGAAACTCTCCCNACNNNCNGT  
GNNCCNTGGGGAAAATGNGNCNNATTCATTTN

#### Ref 11.1

Sequence of BAC4 using primer C2S12, which spans nucleotides 3474-3493 of the cDNA. Exon sequence is not found within this sequence. Since the primer is directed against exon sequence we presume that sequence derived from C2S9 is intron sequence

AGNANNGTTNNGCAGCTGCANNTCTGGACCCANAGGCCGCANGGGCACGAGCCNGGA  
CACGCTCGGCAAAGAGCTGTCCAGAGGGATTCAGAAGCTTCAGGACTGGAAGGGTCTT  
TCGAGCTCAGTTAGCCACCCCCACACCCATTTCAGTTTCACATTTATCTAGTGCTTCCTT  
TTGAATACTTGGGATGTTTTTCTGTTGATCTGTTGGCACTTCCTTCTTCCACAAGACCAG  
AAGCTCATATCCAATCTAAGGTCACTTACCCTTCTGAGAATCTGATGAAAATGGCGTGC  
CTTATGTGCCTAGATGCTTTTGCACACAGTCTAAGGTGACTTATGGACTCCAGGTCCAG  
CAGCCACACCCAGTCCTGGGTCTCCGCACAGGGAGGGACCCGTCTTCACACACCTGTCT  
CAGGTTCTAGCATTGGGCTGCTTCAGCGGTCTCAGGCTGTGAGTAAATGGGATGTGAG  
CTTGGATCGCCCCACGCTGTTGNCCCCCGGGGGGCTTGGCCAGCTGGCCACTTNGAAAT  
GCCTCCTTTTGCCCAGGAAAGCTCACTGCATTTCATGGGGNTTNTCCACGAAGTTCAN  
CTTTANGGG

#### Ref 12.1

Sequence of BAC4 using primer C2S13, which spans nucleotides 3645-3664 of the cDNA. Exon sequence is underlined and represents nucleotides 3683-3699.

AGNAAGGTNNCTCANTNAANNCAGCGTGAGNGTTCAGGTGAGCCAGGCACAGCAGGC  
CGGAGGGCAGCAGGGGACGTCCTTGCCCCTGGGTGACTTGAGAGTCGTTTCCACTAAC  
AAGGTCTACTTGAGAGCCTCGGTTTACCAAGTGATCCCTGCTCCCTTCCCCCAACGTNT  
GTGACATTTCTCCTGATATCAGAGGGGGAGGAAACCTCATGATCCCTGCCCCCGCCCC  
ATGAGGACTGACTGTGGGGACAAAGAGCCAGATCTCATAGACTACCCTGATTTGTCAG  
TATTTGGGGAATTCTGGGTGCCTGATTAGAAGCATCAAGACTCTTCTAAATNCAAAGA  
AGTGTGGAGAGCAGTAGATTTTCCTATAAAACTGGTGTGCTGGTTTCTATGAAAATTG  
GATCCAAAAAAGTCCTTAAGTTTACCCTCTTAATGGNATCTTTTGATTAATGGAATTC  
ATTATTTTAATATAGCCCAATCAATCCAATTTTTCTTTATTGGTAGCATTTTATGTTCTC  
TTTAAAAAAATCTTGGNCTACCTCCAAAATTTACAGATGTTCTCCTAGGGTTTTCTCCTCC  
TTTTGGTTCAAGCATCCCATTCANGTCTTGCAGTCCATTCTGGGG

#### Ref 13.1

Sequence of BAC4 using primer C2S14, which spans nucleotides 4289-4308 of the cDNA. Exon sequence is underlined and represents nucleotides 4321-4448.

GACTTANATTTATTCTTCCTTGCAGAGTAGTGTTAGAATAGATGGCCTACAGAAAAAAA  
AGGTTCTGGGATCTACATGGCAGGGAGGGCTGCACTGACATTGATGCCTGGGGGACCT  
TTTGCCTCGAGGCTGAGCTGGAAAATCTTGAAAATATTTTTTTTTTCTGTGGCACATTC  
AGGTTGAATACAAGAACTATTTTTGTGACTATGTTTTTGATGACCTAAGGGAACTGACC  
ATTGTAATTTTTGTACCANTGAACCANGAGATTTAAGTGCTTTTATATTCATTTCTTGC

Fig. 6A (cont.)

ATTTAAGAAAATATGAAAGCTTAAGGAATTATGTGAGCTTAAAACTAGTCAAGCANTT  
TAGAACCAAAGGCCTATNTTNATAACCGCAACTATGCTNAAAAGNACAAAGTAGTACA  
GNATATTGNTATGTACATATCATTTGGTAATACACNCCNGGCNTTCTGTACATATATGT  
ATTACATTTCTACNTTTTTTAATACTCCCNTGGGCTTATGCCNTTAAGGTAAANTTGNGAT  
AAATTNGGCTGTTCCNGTNTATNCNATAACNCTTTT

**Ref 14.1**

Sequence of BAC4 using primer C2AS15, which spans nucleotides 4680-4700 of the cDNA. Exon sequence is underlined and represents nucleotides 4660-4683.

ATGAGAATGTAATACATATATGTACAGAATGCCAGGACTGTATTAACAATGATATGTA  
CATAACAATATACTGTACTACTTTGTACTTTTCAGCATAGTTGCGGTTATTAATATAGG  
CCTTTGGTTCTAAACTGCTTGACTAGTTTAAAGCTCACATAATTCCTTAAGCTTTCATAT  
TTTCTTAAATGCAAGGAAATGAATATAAAAGCACTAAATCTCCTGGTTCCTGGTACAA  
AAATTACAATGGTCAGTTCCCTTAGGTCATCAAAAAGTAGTCACAAAAAATAGTTCTTGT  
ATTCAACCTGAATGTGCCACAGGAAAAAAAAAATATTTTCAAGATTTTCCAGCTCAGC  
CTCGAGGCAAAAGGCCCCCAAGGCATCAATGTCAGNGCAGCCCTCCTGCCATGTAGATC  
CCAGAACCTTTTTTTTCTGTAGGCCATCTATTCTAACACTACTCTGCAGGGAGAATAAA  
ATCTAAAGNCCAGCTCAAGAGTGCTACCACACCTTTGTTAAGACACAATGAAAACCTTT  
GGATATTGGCAGGNGAGATTTAAAAAAAAAATGTGCCCTTTCTTACCACTCCTATAGNA  
AAGTCTGGTTAAGAAATAACCGTTGGTCTTTATTTTCCTTTTNTTTCCCCTTCCCTTGGG  
NCTTCCTGGGGCTCGG

FIG. 6A (cont.)



HC2A -----  
KIAA ASGNLDKNARFSAIYRQDSNKLSNDDMLKLLADFRKPEKMAKLPVILGNLDITIDNVSSD  
rat -----  
HC4 -----  
HC1 -----  
HC3 -----  
HC5 -----

HC2A -----  
KIAA FPNYVNSSYIPTKQFETCSKTPITFEVEEFVPCIPKHTQPYTIYTNHLYVYPKYLKYDSQ  
rat -----  
HC4 -----  
HC1 -----  
HC3 -----  
HC5 -----

HC2A -----VLHHHQNPETYDEIK  
KIAA KSFAKARNIAICIEFKDSDEEDSQPLKCIYGRPGGPVFTSAFAAVLHHHQNPETYDEIK  
rat -----  
HC4 -----  
HC1 -----  
HC3 -----  
HC5 -----

HC2A IELPTQLHEKHHLTFFHVSCDNSSKGSTKKRDVVETQVGYSWLPLLKDGRVVTSEQHI  
KIAA IELPTQLHEKHHLTFFHVSCDNSSKGSTKKRDVVETQVGYSWLPLLKDGRVVTSEQHI  
rat -----  
HC4 -----  
HC1 -----  
HC3 -----  
HC5 -----

HC2A PVSANLPSGYLGYQELGMGRHYGPEIKWVDGGKPLLKISTHLVSTVYTQDQHLHNFFQYC  
KIAA PVSANLPSGYLGYQELGMGRHYGPEIKWVDGGKPLLKISTHLVSTVYTQDQHLHNFFQYC  
rat -----  
HC4 -----  
HC1 -----  
HC3 -----GPGPARSTVSIISLISNSARV  
HC5 -----

HC2A QKTESGAQALGNELVKYLKSLHAMEGHVMIAFLPTILNQLFRVLT-RATQEEVAVNVTRV  
KIAA QKTESGAQALGNELVKYLKSLHAMEGHVMIAFLPTILNQLFRVLT-RATQEEVAVNVTRV  
rat -----  
HC4 -----MEIQVLIRFLSVILMQLFWLPMIHEDDVPISCPMV  
HC1 -----MSFLPIILNQLFKVLV-QNEEDEITTTVTRV  
HC3 NRSRSLSNSNPDISGTPTSPDDEVRSIIGSKGLDRSNSWNTGGPKAAPWGSNPSPSAES  
HC5 -----

Fig. 6B(cont.)

HC2A I I H V V A Q C H E E C L E S H L R S Y V K Y A Y K A E P Y V A S E Y K T V H E E L T R S M T T I L K P S A D F L T S N  
 KIAA I I H V V A Q C H E E G L E S H L R S Y V K Y A Y K A E P Y V A S E Y K T V H E E L T K S M T T I L K P S A D F L T S N  
 rat -----  
 HC4 L F H I V S K C H E E G L D S Y L S S F I K Y S F R P G K P S A P Q A P L I H E T L A T M M I A L L K Q S A D F L A I N  
 HC1 L P D I V A K C H E E Q L D H S V Q S Y I K F V F K T R --- A C K E R P V H E D L A K N V T G L L K - S N D S P T V K  
 HC3 T Q A M D R S C N R M S S H T E T S S F L Q T L T G R L P --- T K K L F H E E L A L Q W V V C S G --- S V R --- E  
 HC5 -----

Cadherin  
 Cleavage

HC2A K L L R Y S W F F F D V L I K S M A Q H L I E N S K V K L I R N Q R F P A S Y H H A A E T V V N M L M P H I T Q K F G D  
 KIAA K L L K Y S W F F F D V L I K S M A Q H L I E N S K V K L I R N Q R F P A S Y H H A V E T V V N M L M P H I T Q K F R D  
 rat -----  
 HC4 K L L K Y S W F F F E I I A K S M A T Y L L E E N K I K L T H G Q R F P K A Y H H A L H S L F L A I T - I V E S Q Y A E  
 HC1 H V L K H S W F F F A I I L K S M A Q H L I D T N K I Q L E R P Q R F P E S Y Q N E L D N L V M V L S D H V I W K Y K D  
 HC3 S A L Q Q A W F F F E L M V K S M V H H L Y F N D K L E A P R K S R F P E R F M D D I A A L V S T I A S D I V S R F Q K  
 HC5 -----

6.1.  
 1.1 / 1.2 / 2.1 / 2.2

HC2A N P E A S K N A N H S L A V F I K R C F T F M D R G F V F K Q I N --- N Y I S --- C F A P G D P K T L F E Y K F E F L 2.1  
 KIAA N P E A S K N A N H S L A V F I K R C F T F M D R G F V F K Q I N --- N Y I S --- C F A P G D P K T L F E Y K F E F L  
 rat -----  
 HC4 I P K E S R N V N Y S L A S F L K C C L T L M D R G F V F N L I N --- D Y I S --- G F S P K D P K V L A E Y K F E F L  
 HC1 A L E E T R R A T H S V A R F L K R C F T F M D R G C V F K M V N --- N Y I S --- M F S S G D L K T L C Q Y K F D F L 7.1  
 HC3 D T E M V E R L N T S L A F F L N D L L S V M D R G F V F S L I K S C Y Q V S S K L Y S L P N P S V I V S L R L D F L 3.1 / 3.2  
 HC5 -----

HC2A R V V C N H E H Y I P L N L P M --- P F G K G R I Q R --- Y Q D L Q L --- D Y S L T D E F  
 KIAA R V V C N H E H Y I P L N L P M --- P F G K G R I Q R --- Y Q D L Q L --- D Y S L T D E F  
 rat -----  
 HC4 Q T I C N H E H Y I P L N L P M --- A F A K P K L Q R --- V Q D S N L --- E Y S L S D E Y  
 HC1 Q E V C Q H E H F I P L C L P I R S A N I P D P L T P S E S --- T Q E L H A S D M P E Y S V T N E F  
 HC3 R I I C S H E H Y V T L N L P C S L L T P P A S P S P S V S S A T S Q S S G F S T N V Q D Q K I A N M F E L S --- V P F 4.1 / 4.2  
 HC5 ----- M N A D T A P T S P C P S I S --- S Q N S S C S S F Q D Q K I A S M F D R T S R V P A

HC2A C R N H F L V G L L L R E V G T A L Q E F R E --- V R L I A I S V L K N L L I K H S F D D R Y A S R S H Q A R I A T 3.1  
 KIAA C R N H F L V G L L L R E V G T A L Q E F R E --- V R L I A I S V L K N L L I K H S F D D R Y A S R S H Q A R I A T  
 rat -----  
 HC4 C K H H F L V G L L L R E T S I A L Q D N Y E --- I R Y T A I S V I K N L L I K H A F D T R Y Q H K N Q Q A K I A Q  
 HC1 C R K H F L I G I L L R E V G F A L Q E D Q D --- V R H L A V L K N L M A K H S F D D R Y R E P R K Q A Q I A S 8.1  
 HC3 R Q Q H Y L A G L V L T E L A V I L D P D A E G L F G L H K K V I N M V H N L L S S H D S D P R Y S D P Q I K A R V A M  
 HC5 S S T S - S P G L L F T E L A A A L D A E G E G I S E V Q R K A V S A I H S L L S S H D L D P R C V K P E V K V K I A A

HC2A L Y L P L F G L L I E N V Q R I N V R D V S P F P V N A G - M T V K D E S L A L P A V N P L V T P Q K G S T L D N S L H  
 KIAA L Y L P L F G L L I E N V Q R I N V R D V S P F P V N A G - M T V K D E S L A L P A V N P L V T P Q K G S T L D N S L H  
 rat -----  
 HC4 L Y L P F V G L L L E N I Q R L A G R D T L Y S C A A M P N S A S R D E F P C G --- F T S P --- A N --- R G S L S  
 HC1 L Y M P L Y G M L L D N M P R I Y L K D L Y P F T V N T S N Q G S R D D L S T N G G F Q S Q T A I K H A N S V D T S F S 9.1  
 HC3 L Y L P L I G I I M E T V P Q L Y D F T E T H N Q R G R P I C I A T D D Y E S E --- S G --- S M I S  
 HC5 L Y L P L V G I I L D A L P Q L C D F T V A D T R R Y R --- T S G S D E E Q E --- G A --- G A I T

4.1 / 4.2

HC2A K D L L G A I S G I A S P Y T T S T P N I N S V R N A D S R G S L I S T D S G N S L P E R N S E K S N S L D K H Q Q S S 5.1 / 5.2  
 KIAA K D L L G A I S G I A S P Y T T S T P N I N S V R N A D S R G S L I S T D S G N S L P E R N S E K S N S L D K H Q Q S S  
 rat -----  
 HC4 T D K D T A Y G S F Q N G --- H G I K R E D S R G S L I P - E G A T G F P D Q G N T G E N --- T R Q S  
 HC1 K D V L N S I A F S S --- I A I S T V N H A D S R A S L A S L D S N P S T N E K S S E K T D N C E K I P R P L 10.1  
 HC3 Q T V A M A I A G T S V P Q --- L T R P G S F L L T S T S G R Q H T --- 3.1  
 HC5 Q N V A L A I A G N N F N --- L K T S G - I V L S S L P Y K Q Y N --- 2.1

HC2A	TLGNSVVRCDKLDQSEIKSLLMCFLYILKSMSDDALFTYWN-KASTSELMDFFTISEVCL	
KIAA	TLGNSVVRCDKLDQSEIKSLLMCFLYILKSMSDDALFTYWN-KASTSELMDFFTISEVCL	
rat	-----	
HC4	STRSSVSQYNRLDQYEIRSLLMCYLYIVKMISEDTLLTYWN-KVSPQELINILILLEVCL	
HC1	ALIGSTLRFDRLDQAETRSLLMCFHIMKTISYETLIAYWQ-RAPSPEVSDFFSIIPVCL	11.1 / 11.2
HC3	-----TFSAESSRSLICLLWVLKN-ADETVLQKWFTDLSVLQNLRLDLLYLVCV	
HC5	-----MLNADTTRNLMICFLWIMKN-ADQSLIRKWIADLPSTQLNRILDLLFICV	
HC2A	HQFQYMGKRYIARNQEGLG--PIVHDRKS-----QTLPVSRNRTGMM	6.1
KIAA	HQFQYMGKRYIAR-----TGMM	
rat	-----	
HC4	FHFRYMGKRNIARVHDAWLSKHFGIDRKS-----QTMPALNRNRSGVM	
HC1	QNFRLGKRNIIRKIAAAF--KFVQSTQNNGTLKGSNPSCQTSGLLAQWMHSTSRHEGHK	
HC3	SCFEYKGGKVFERMNSLTFK--KSKDMRAK-----LEEAILGSIGARQEMV	
HC5	LCFEYKGGKQSSDKVSTQVLQ--KSRDVKAR-----LEEALLRGEGARGEMM	
HC2A	HARLQQL-----GSLDNS-----LTFNHSYGHSDADVLHQSLLEANIATEVC	
KIAA	HARLQQL-----GSLDNS-----LTFNHSYGHSDADVLHQSLLEANIATEVC	
rat	-----	
HC4	QARLQHL-----SSLESS-----FTLNHSSTTTEADIFHQALLEGNTATEVS	
HC1	QHRSQTLPIIRGK--NALSNPKL----LQMLDNTMTSNEIDIVHHVDTEANIATEGC	12.1 / 12.2
HC3	RRSRGQLERSPSGSAFGSQENLRWRKDMTHWRQNTTEKLDKSRAEIEHEALIDGNLATEAN	6.1 / 6.2
HC5	RRRAPGNDRFP-----GLNENLRWKKEQTHWRQANEKLDKTKAELDQREALISGNLATEAH	
HC2A	LTALDTLSLFTLAFKNQLLADHGHNPLMKKVFDVYLCFLQKHQSETALKNVFTALRSLIY	7.1
KIAA	LTALDTLSLFTLAFKNQLLADHGHNPLMKKVFDVYLCFLQKHQSETALKNVFTALRSLIY	
rat	-----KLSRGHSPLMKKVFDVYLCFLQKHQSEMALKNVFTALRSLIY	
HC4	LTVLDTISFFTQCFKTHFLNNDGHNPLMKKVFDIHLAFLKNGQSEVSLKHVFASLRAFIS	
HC1	LTILDVLSLFTQTHORQLOQCDCONSLMKRGFDTYMLFFQVNOQATALKHVFASLRLFVC	13.1
HC3	LIILDTLEIVVQTVS--VTES--KESILGGVLKVLHSMACNQSAVYLQHCFAQORALVS	
HC5	LIILDMQENIIQASS--ALDC--KDSLLGGVLRVLVNSLNCQSTTYLTHCFATLRALIA	3.1
HC2A	KFPSTFYEGRADMCAALCYEILKCCNSKLSSIRTEASQLLYFLMRNDFDYTGKKSFVRTH	
KIAA	KFPSTFYEGRADMCAALCYEILKCCNSKLSSIRTEASQLLYFLMRNDFDYTGKKSFVRTH	
rat	KFPSTFYEGRADMCASLCYEVLKCCNSKLSSIRTEASQLLYFLMRNDFDYTGKKSFVRTH	
HC4	KFPSAFFKGRVNMCAAFCEVLKCCCTSKISSTRNEASALLYLLMRNDFEYTKRKTFLRTH	
HC1	KFPSAFFQGPADLCGSFCYEVLKCCNHRSRSTQTEASALLYLFMRKNFEFNKQKSIVRSH	
HC3	KFPPELLFEEETEQCADLCLRLLRHCSSSIGTIRSHPSASLYLLMRQNFIGN--NFARVK	7.1 / 7.2
HC5	KFGDLLFEEVEQCDFDLCHQVLHHCSSSMDVTRSQCACATLYLLMRFSFGATS--NFARVK	
HC2A	LQVIIISVSQLIADVVGIGETRFOQSLSIINNCANSRDLIKHTSFSSDVKDLTKRIRTVLM	
KIAA	LQVIIISVSQLIADVVGIGETRFOQSLSIINNCANSRDLIKHTSFSSDVKDLTKRIRTVLM	
rat	LQVIIISLSQLIADVVGIGETRFOQSLSIINNCANSRDLIKHTSFSSDVKDLTKRIRTVLM	
HC4	LQIIIAVSQLIADVALSGGSRFQESLFIINNANSRDRPMLARAFPAEVKDLTKRIRTVLM	
HC1	LQLIKAVSQLIAD-AGIGGSRFQHS LAITNNFANGDKOMKSNFPAEVKDLTKRIRTVLM	14.1 / 14.2 / 15.1 / 15
HC3	MQVPMSLSSLVGTSQNFNEEFLRRSLKTIITYAEEDLELRETTFPDQVQDLVFNLMILS	
HC5	MQVTMSLASLVGRAPDFNEEHLRRSLRTILAYSEEDTAMQMTFPPTQVEELLCNLNSILY	

HC2A	ATAQMKEHENDPEMLVDLQYSLAKSYASTPELRKTWLD SMARIHVKN GDLSEAAMCYVHV	
KIAA	ATAQMKEHENDPEMLVDLQYSLAKSYASTPELRKTWLD SMARIHVKN GDLSEAAMCYVHV	
rat	ATAQMKEHENDPEMLVDLQYSLAKSYASTPELRKTWLD SMARIHVKN GDLSEAAMCYVHV	
HC4	ATAQMKEHEKDPEMLIDLQYSLAKSYASTPELRKTWLD SMAKIHVKN GDLSEAAMCYVHV	
HC1	ATAQMKEHEKDPEMLVDLQYSLANSYASTPELRRTWLE SMAKIHARNGDLSEAAMCYIHI	16.1 / 16.2
HC3	DTVKMKEHQEDPEMLIDL MYRIAKGYQTSPDLRLTWLQNMAGKHSERSNHAEEAAQCLVHS	
HC5	DTVKMREFQEDPEMLMDL MYRIAKSYQASPDRLRLTWLQNMAEKHTKKKCYTEAAMCLVHA	

		SH3	
HC2A	TALVAEYI	TRKGV-----	-----FRQGCTAFRVITPN
KIAA	TALVAEYI	TRKEA-----	-----VQWEPPLLPHSHSACLRRSRGGVFRQGCTAFRVITPN
rat	TALVAEYI	TRKEAD-----	-----LALQREPPVFPYSHTSCQRKSRGGMFRQGCTAFRVITPN
HC4	AALVAEFT	HRKKL-----	-----FPNGCSAFKKITPN
HC1	AALIAEYI	KRKG YWKVEKICTASLLSEDT HPCDSNSLLTTPSGGSMFSGMGP AFLSITPN	
HC3	AALVAEYI	SMLED-----	-----RKYLPVGC VTFQNISSN
HC5	AALVAEYI	SMLED-----	-----HSYLPVGSV SFQNISSN

HC2A	IDEEASMMEDVGMQD-----	VHFNE DVLME LLEQCADGLWKAERYELIADIYKLI IPI	9.1
KIAA	IDEEASMMEDVGMQD-----	VHFNE DVLME LLEQCADGLWKAERYELIADIYKLI IPI	
rat	IDEEASMMEDVGMQD-----	VHFNE DVLME LLEQCADGLWKAERLRAGLLTSINSSSP	
HC4	IDEEGAMKEDAGMMD-----	VHYSEEVLLELLEQCVNGLWKAERYEI ISEISKLI GPI	
HC1	IKKEGAAKEDSGMHD-----	TPYNE NILVEQLYMCGEFLWK SERYELIADV NKP IIAV	17.1 / 17.2
HC3	VLEESAVSDDV VSPDEEGICSGKYFTESGLVGLLEQAAASF SMAGMYEAVNEVYKVL IPI		
HC5	VLEESV VSED TLSPDEDGVCAGQYFTESGLVGLLEQAAELFSTGGLYETVNEVYKVL IPI		

		ITAM	ITAM	ITAM	ITAM	
HC2A	YEKRRD-----					9.1
KIAA	YEKRRD FERLAHL YDTLH RAYSKVTEVMHSGRLLGTYFRVAFFGQAAQYQFTDSETDVE					
rat	SMKSGGTLETHLYDTLH RAYSKVTEVITR-----					
HC4	YENRREFENLTQVYRTLHGAYTKILEVMHTKKRLLG-----					
HC1	FEKQRDFKKLS DLYYDIHRSY LKVAEVVNSEKRLFG-----					
HC3	HEANRDAKKLSTIHGKLQEA FSKIVHQSTGWERMFG-----					9.1
HC5	LEAHREFRKLTLTHSKLQRAFDSIYNKDH--KRMFG-----					

HC2A	-FFED EDGKEYIYKEPKLTPLSEISQRLLKLYSDKFGSENVKMIQDSGKVNPKDLDSKYA	10.1
KIAA	GFFED EDGKEYIYKEPKLTPLSEISQRLLKLYSDKFGSENVKMIQDSGKVNPKDLDSKYA	
rat	GFFED EDGKEYIYKEPKLTPLSEISQRLLKLYSDKFGSENVKMIQDSGKVNPKDLDSKFA	
HC4	SFFEE EDGKEYIYKEPKLTGLSEISLRLVKLYGEKFGTENVKIIQSDKVNAKELDPKYA	
HC1	GFFEE EEGKEYIYKEPKLTGLSEISQRLLKLYADKFGADNVKIIQDSNKNPNKDLDPKYA	
HC3	TKFGDLDEQEFVYKEPAITKLAEISHRLEGFYGERFGEDVVEVIKDSNPVDKCKLDPNKA	10.1 / 10.2
HC5	SKFGDLDEQEFVYKEPAITKLPEISHRLEAFYGCFGAEFVEVIK DSTPVDKTKLDPNKA	4.1

HC2A	YIQVTHVIPFFDEKELQERKTEFERSHNIRRFMFEMPFTQTGKRQGGVEEQCKRRTILTA	11.1 / 11.2
KIAA	YIQVTHVIPFFDEKELQERKTEFERSHNIRRFMFEMPFTQTGKRQGGVEEQCKRRTILTA	
rat	YIQVTHVTPFFDEKELQERKTEFERCHNIRRFMFEMPFTQTGKRQGGVEEQCKRRTILTA	
HC4	HIQVTYVKPYFDDKELTERKTEFERNHNISR FVFEAPYTL SGKKQGCIEEQCKRRTILT	
HC1	YIQVTYVTPFFEEKEIEDRKTDFEMHHNINRFVFETPFTLSGKKHGGVAEQCKRRTILT	18.1
HC3	YIQITYVEPYFDTYEMKDRITYFDKNYNLRRFMYCTPFTLDGRAHGE LHEQFKRRTILT	
HC5	YIQITFVEPYFDEYEMKDRVTYFEKNFNLRRFMYTTPFTLEGRPRGELHEQYRRNTVLT	

FIG. 6B (cont.)

Coiled-Coil 1

HC2A	IHC	FPYVKKRIPV	MYQHHTDLN	IEVAIDEMSKKVAELRQLC	SSAEVDMIKLQLKLOGSV
KIAA	IHC	FPYVKKRIPV	MYQHHTDLN	IEVAIDEMSKKVAELRQLC	SSAEVDMIKLQLKLOGSV
rat	IHC	FPYVKKRIPV	MYQHHTDLN	IEVAIDEMSKKVAELHQLC	SSAEVDMIKLQLKLOGSV
HC4	SNS	FPYVKKRIPIN	CEQQINLKE	IDGATDEIKDKTAELOKLC	SSTDVDMIQLQLKLOGSV
HC1	SHL	FPYVKKRIQV	ISQSSTELN	IEVAIDEMSRKVSELNQLCT	MEEVDMISLQLKLOGSV
HC3	SHAF	PIKTRVNVTH	KEIILTH	IEVAIEDMQKKTOELAFATH	QDPADPKMLQMVLOGSV
HC5	MHAF	PIKTRISVIQ	KEEFVLTH	IEVAIEDMKKKTLQLAVAIN	QEPDAKMLQMVLOGSV

11.1

Coiled-Coil 2

HC2A	SVQ	VNAGPLAYARA	FLDDTNTKRY	PDNKVKLLKEVFRQFVEAC	QALAVNERLIKEDQLE
KIAA	SVQ	VNAGPLAYARA	FLDDTNTKRY	PDNKVKLLKEVFRQFVEAC	QALAVNERLIKEDQLE
rat	SVQ	VNAGPLAYARA	FLDDTNTKRY	PDNKVKLLKEVFRQFVEAC	QALAVNERLIKEDQLE
HC4	SVQ	VNAGPLAYARA	FLNDSQASKY	PPKKVSELKDMFRKFIOAC	SIALELNERLIKEDQVE
HC1	SVK	VNAGPMAYARA	FLLEETNAKKY	PDNQVKLLKEIFRQFADAC	QALDVNERLIKEDQLE
HC3	GTT	VNOGPLEVAQ	VFLSEIPSDPK	LFRHHNKLRLCFKDFTKR	CEDALRKNKSLIGPVQKE
HC5	GAT	VNOGPLEVAQ	VFLAEIPADPK	LYRHHNKLRLCFKEFIMRC	GEAVEKNKRLITADQRE

11.1 / 12.1

Coiled-Coil 2

HC2A	YQE	EMKANYREMA	KELSEIMHEQ	ICPLEEKTS-VLPNSLHI	FNAISGTPTSTMVHGMTSS
KIAA	YQE	EMKANYREMA	KELSEIMHEQ	LG-----	
rat	YQE	EMKANYREIR	KELSDIIVPR	ICPGEDKRATKFP	PAHLQRHQRTDNKHSGSRVDQFILS
HC4	YHE	GLKSNFRDM	VKELSDIIEQ	ILQEDTMHSPWMSNTLHV	FCAISGTSSDRGYGSPRYA
HC1	YQE	ELRSHYKDM	LSELSTVMNE	QITGRDDLSK---RGVDQ	TCTRVISKATPALPTVSISS
HC3	YQRE	LG----	KLSS-----	PZ-----	
HC5	YQO	ELKKNYNKL	KENLRPMIER	KIPELYKPIFRVESQKRDS	FHRSSFRKCETQLSQGSZ-

19.1

PBM

HC2A	SSVVZ	-----
KIAA	-----	
rat	CVTL	PHEPHVGTCFVMCKLRTTFRANHWFCQAQEEAMNGREKEPWTVI
HC4	EVZ	-----
HC1	SAEVZ	-----
HC3	-----	
HC5	-----	

HC2A	-----
KIAA	-----
rat	VHIFF
HC4	-----
HC1	-----
HC3	-----
HC5	-----



**HindIII**  
**Eco RI**

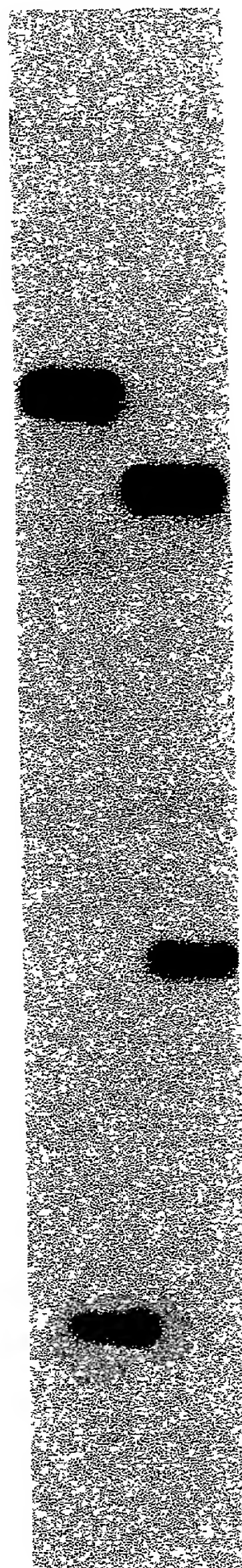


← ~ 4.5 kb

← ~ 1.85 kb

genomic DNA

**Pst / Eco RI**



← ~ 4.5 kb

← ~ 1.85 kb

## BAC 6 DNA

FIG. 7

00687837 101300

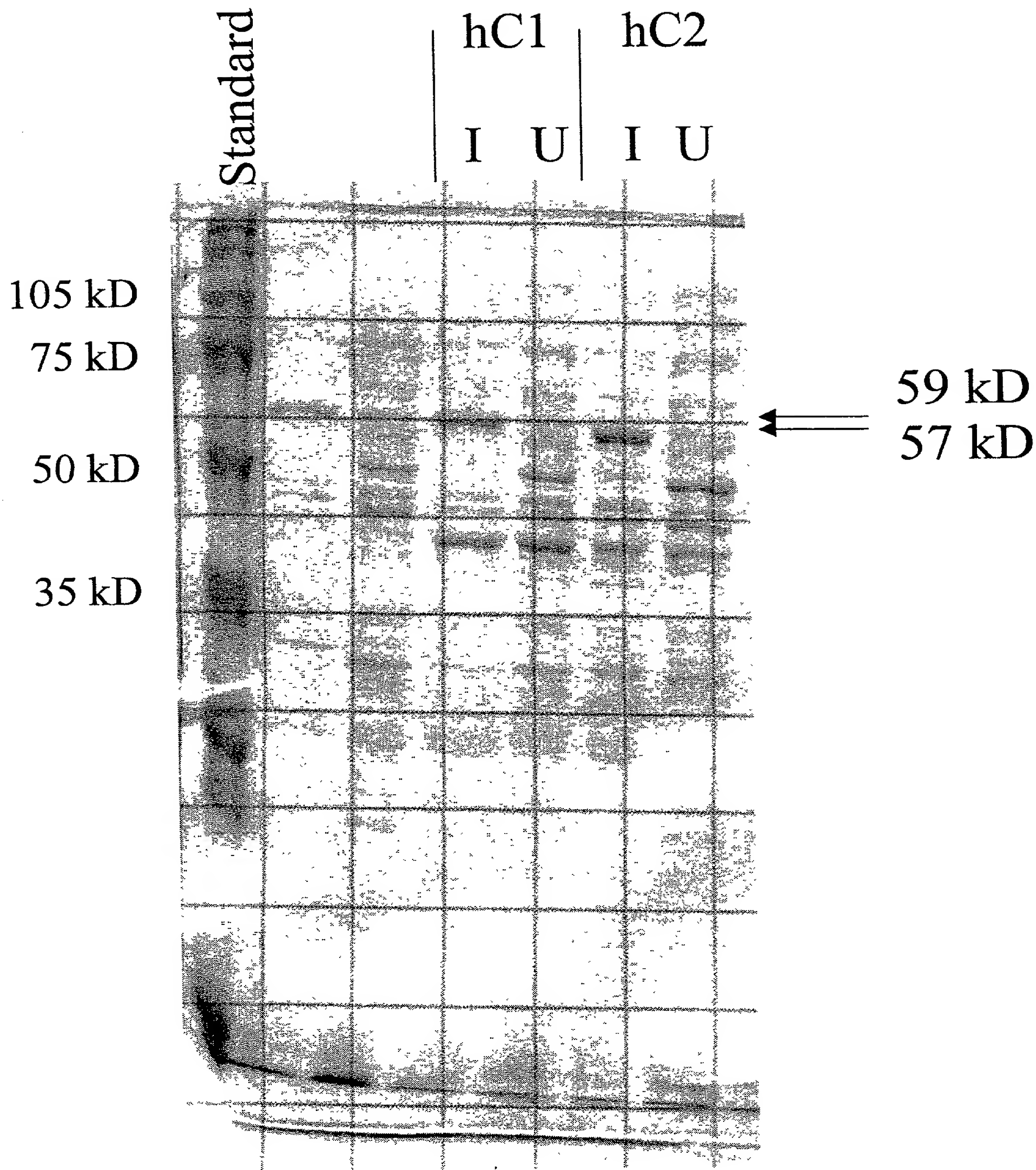


FIG. 8

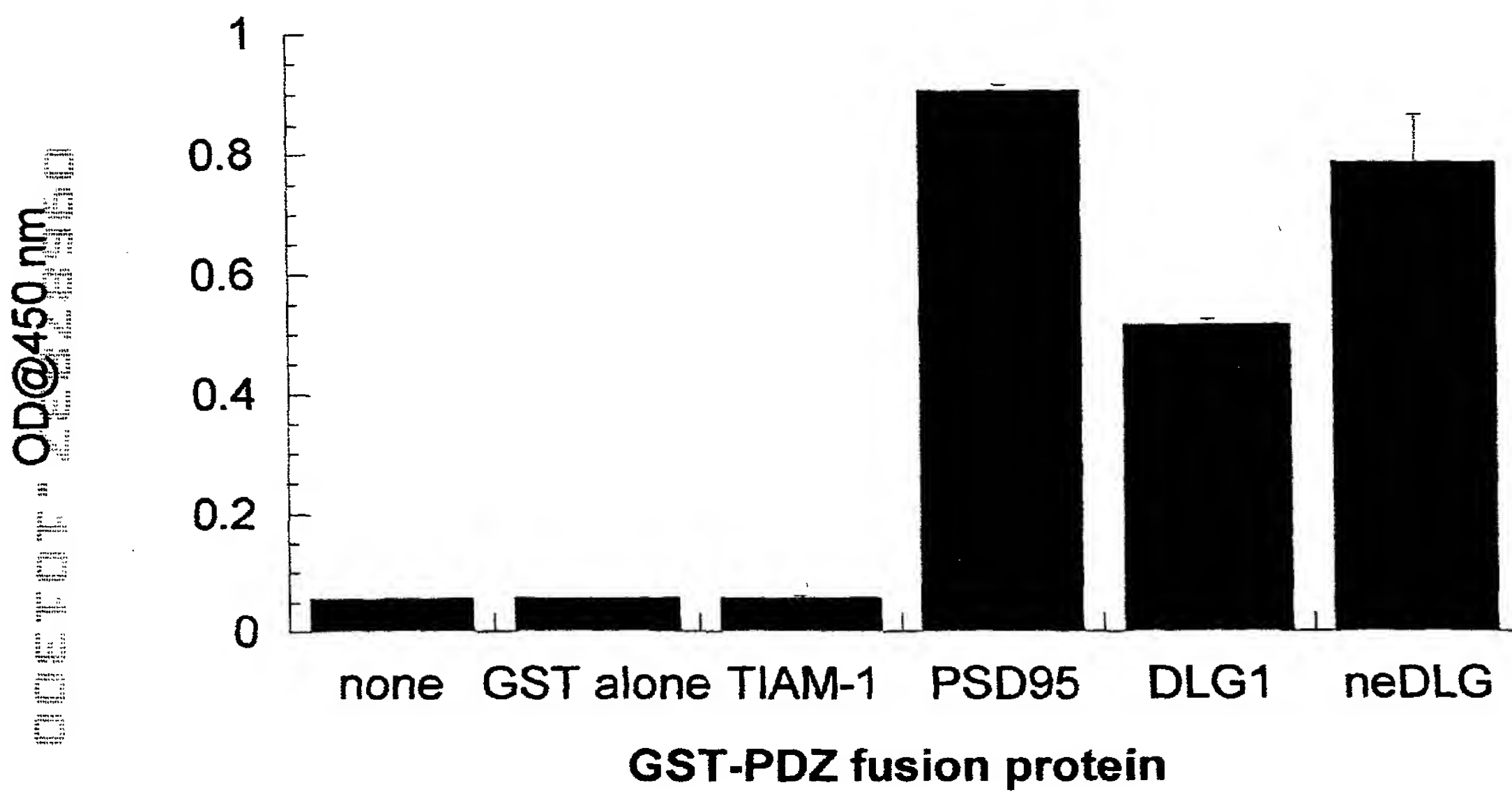


FIG. 9A

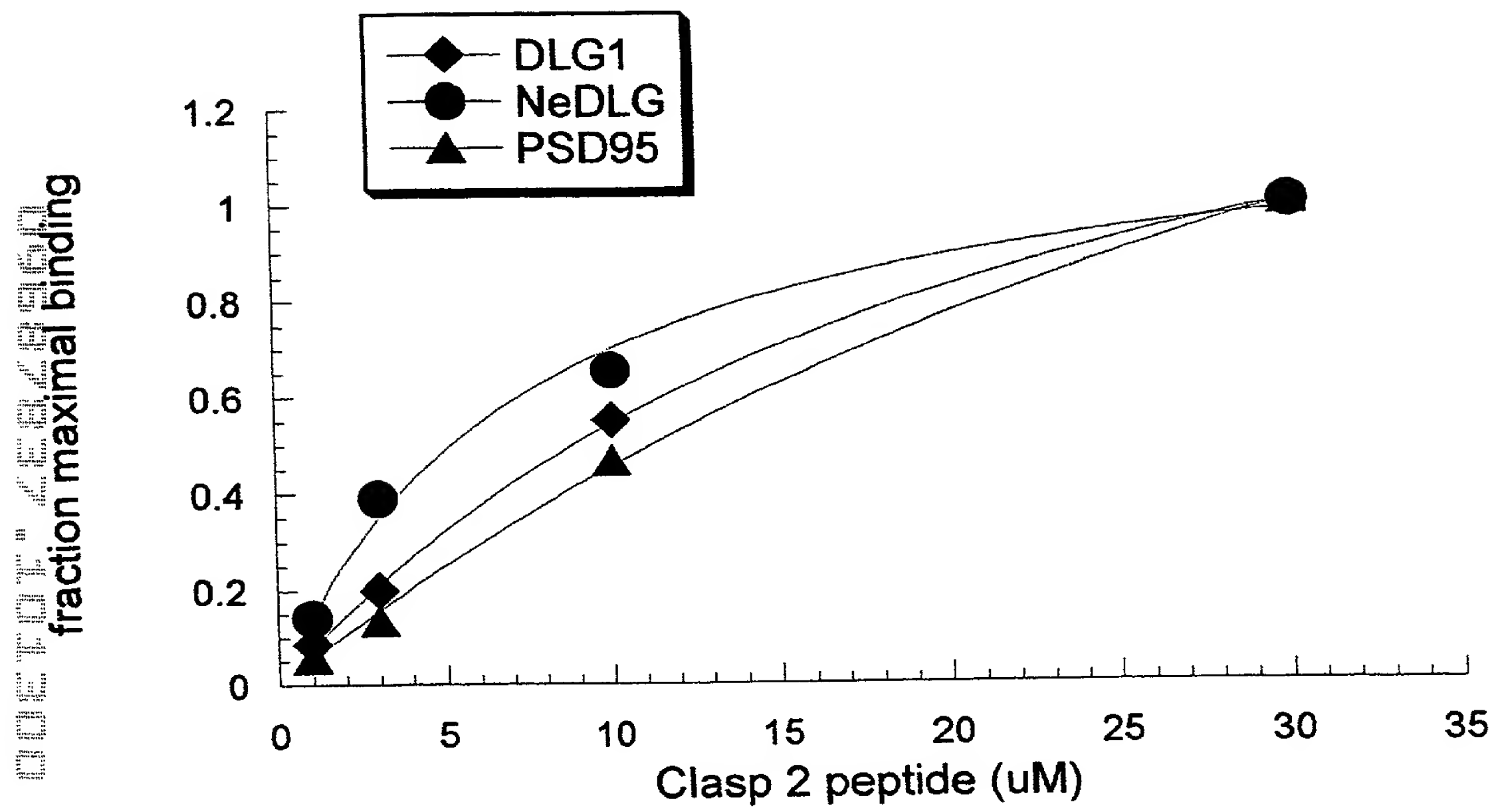


FIG. 9B

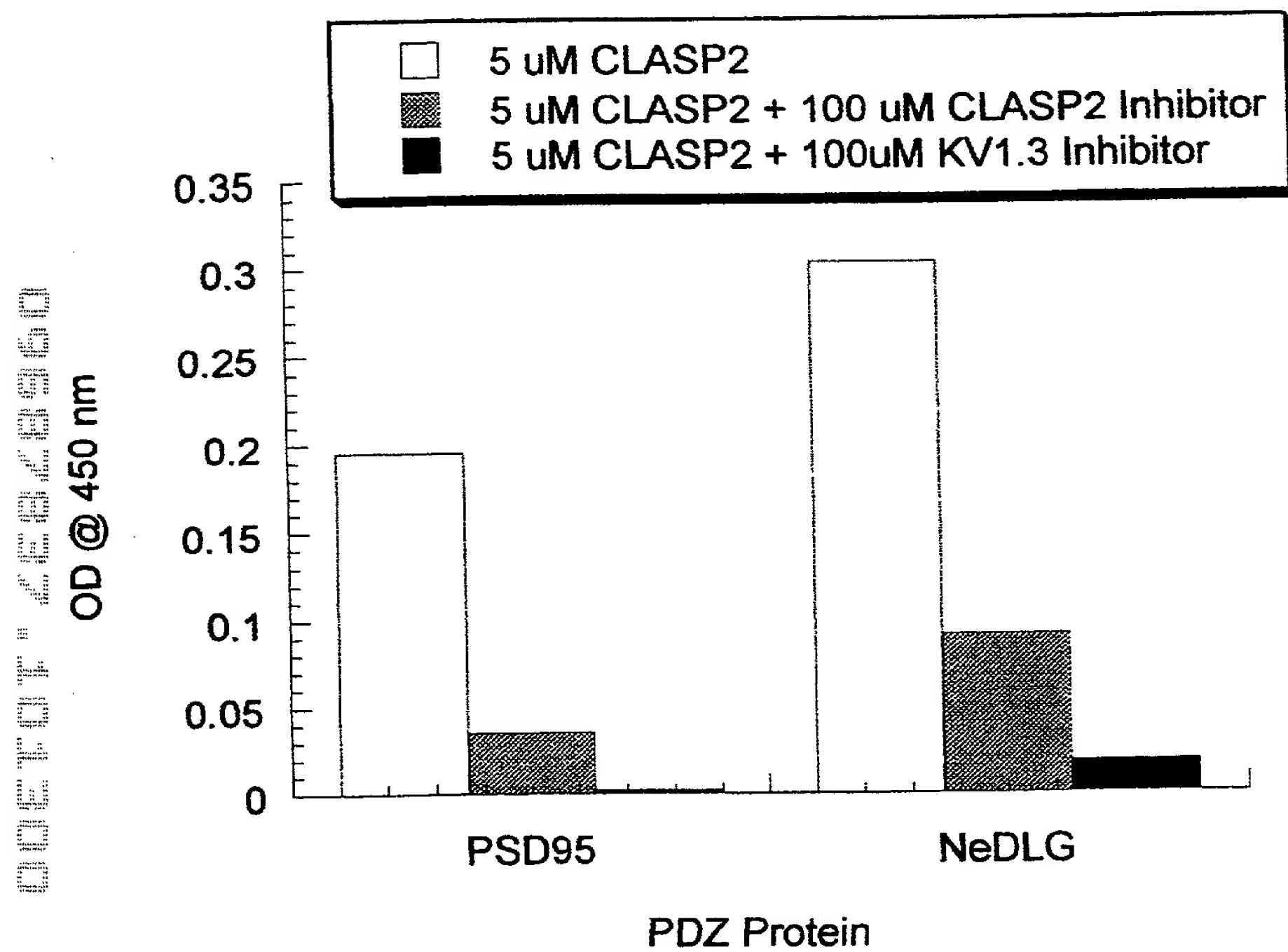


Fig. 9C



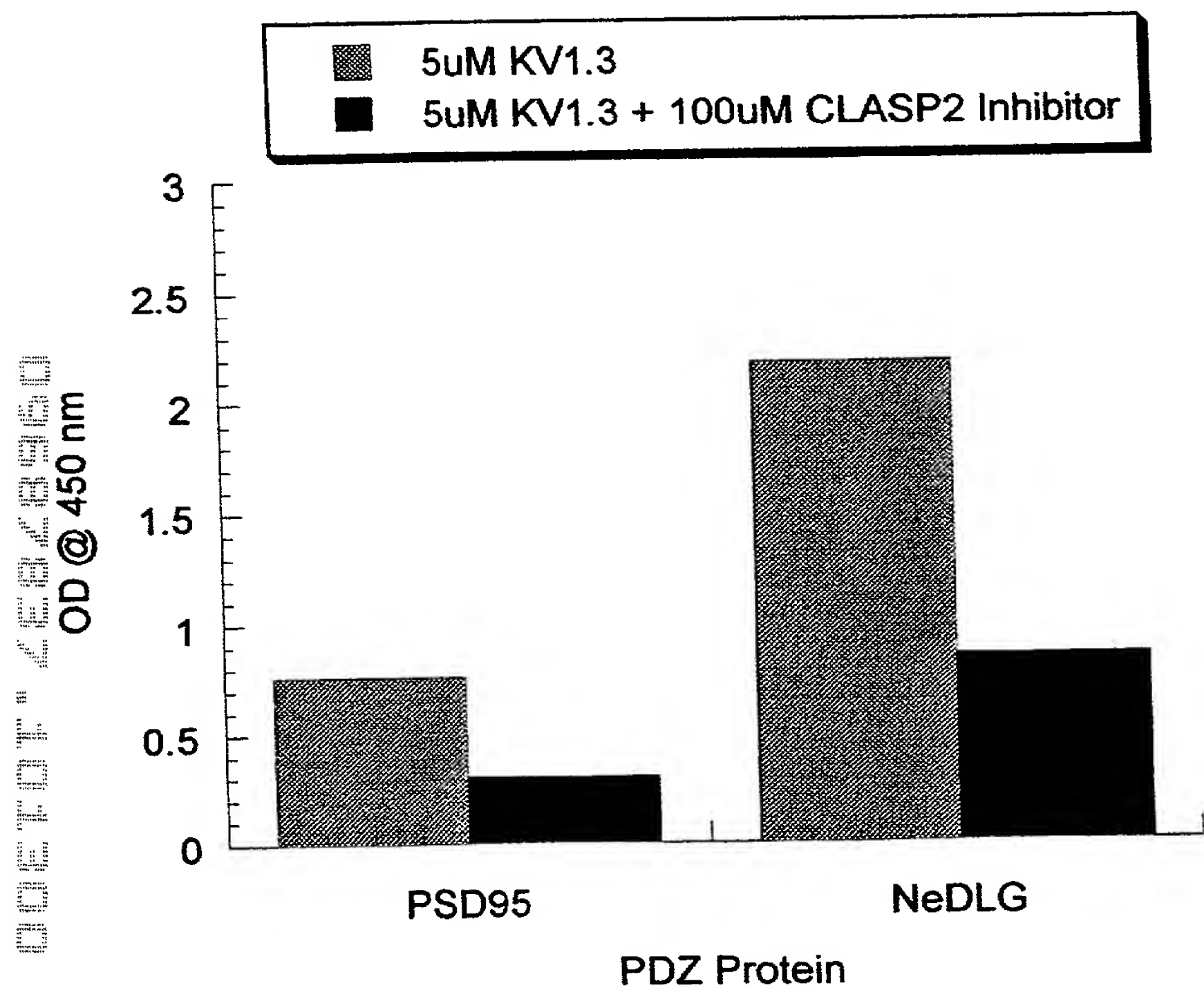


FIG. 9D

	10	20	30	40	50	60	70	80
1	AATTGTAATA	CGACTCACTA	TAGGGCGAAT	TGGGTACCGG	GCCCCCCTC	GAGGTGACG	GTATCGATAA	GCTTGATATC 80
81	GAATTCGGCA	CGAGTTTTAC	ACCATCACCA	AAACCCAGAA	TTTTATGATG	AGATTAAAAT	AGAGTTGCCC	ACTCAGCTGC 160
161	ATGAAAAGCA	CCACCTGTTG	CTCACATTCT	TCCATGTCAG	CTGTGACAA	TCAAGTAAAG	GAAGCACGAA	GAAGAGGGAT 240
241	GTCGTTGAAA	CCCAAGTTGG	CTACTCCTGG	CTTCCCCTCC	TGAAAGACGG	AAGGCTGGTG	ACAAGCGAGC	AGCACATCCC 320
321	GGTCTCGGCG	AACCTTCCTT	CGGGCTATCT	TGGCTACCAA	GAGCTTGGGA	TGGGCAGGCA	TTATGGTCCG	GAAATTAAAT 400
401	GGGTAGATGG	AGGCAAGCCA	CTGCTGAAAA	TTTCCACTCA	TCTGGTTTCT	ACAGGGATAC	TCAGGATCAG	CATTTACATA 480
481	ATTTTTTCCA	GTACTGTCAG	AAAACCGAAT	CTGGAGCCCC	AGCCTTAGGA	AACGAACTTG	TAAAGTACCT	TAAGAGTCTG 560
561	CATGCGATGG	AAGGCCACGT	GATGATCGCC	TTCTTGCCCC	CTATCCTAAA	CCAGCTGTTT	CGAGTCCTCA	CCAGAGCCAC 640
641	ACAGGAAGAA	GTCGCGGTTA	ACGTGACTCG	GGTCATTATT	CATGTGGTTG	CCCAGTGCCA	TGAGGAAGGA	TTGGAGAGCC 720
721	ACTTGAGGTC	ATATGTTAAG	TACGCGTATA	AGGCTGAGCC	ATATGTTGCC	TCTGAATACA	AGACAGTGCA	TGAAGAACTG 800
801	ACCAAATCCA	TGACCACGAT	TCTCAAGCCT	TCTGCCGATT	TCCTCACCAG	CAACAACTA	CTGAGGTACT	CATGGTTTTT 880
881	CTTTGATGTA	CTGATCAAA	CTATGGCTCA	GCAATTTGATA	GAGAATCCCA	AAGTTAAGTT	GCTGCGAAAC	CAGAGATTTC 960
961	CTGCATCCTA	TCATCATGCA	GCGGAAACCG	TGTAAATAT	GCTGATGCCA	CACATCACTC	AGAAGTTTGG	AGATAATCCA 1040
1041	GAGGCATCTA	AGAACGCGAA	TCATAGCCTT	GCTGTCTTCA	TCAAGAGATG	TTTCACCTTC	ATGGACAGGG	GCTTTGTCTT 1120
1121	CAAGCAGATC	AACAACCTACA	TTAGCTGTTT	TGCTCCTGGA	GACCCAAAGA	CCCTCTTTGA	ATACAAGTTT	GAATTTCTCC 1200
1201	GTGTAGTGTG	CAACCATGAA	CATTATATTC	CGTTGAACTT	ACCAATGCCA	TTTGGAAAAG	GCAGGATTCA	AAGATACCAA 1280
1281	GACCTCCAGC	TTGACTACTC	ATTAACAGAT	GAGTTCTGCA	GAAACCACTT	CTTGGTGGGA	CTGTTACTGA	GGGAGGTGGG 1360
1361	GACAGCCCTC	CAGGAGTTCC	GGGAGGTCCG	TCTGATCGCC	ATCAGTGTGC	TCAAGAACCT	GCTGATAAAG	CATTCTTTTG 1440
1441	ATGACAGATA	TGCTTCAAGG	AGCCATCAGG	CAAGGATAGC	CACCTCTAC	CTGCCTCTGT	TTGGTCTGCT	GATTGAAAAC 1520
1521	GTCCAGCGGA	TCAATGTGAG	GGATGTGTCA	CCCTTCCCTG	TGAACGCGGG	CATGACCGTG	AAGGATGAAT	CCCTGGCTCT 1600
1601	ACCAGCTGTG	AATCCGCTGG	TGACGCCGCA	GAAGGGAAGC	ACCCTGGACA	ACAGCCTGCA	CAAGGACCTG	CTGGGCGCCA 1680
1681	TCTCCGGCAT	TGCTTCTCCA	TATACAACCT	CAACTCCAAA	CATCAACAGT	GTGAGAAATG	CTGATTCGAG	AGGATCTCTC 1760
1761	ATAAGCACAG	ATTCGGGTAA	CAGCCTTCCA	GAAAGGAATA	GTGAGAAGAG	CAATTCCTTG	GATAAGCACC	AACAAAGTAG 1840
1841	CACATTGGGA	AATTCCGTGG	TTCCGTGTGA	TAAACTTGAC	CAGTCTGAGA	TTAAGAGCCT	ACTGATGTGT	TTCTCTTACA 1920
1921	TCTTAAAGAG	CATGCTGTAT	GATGCTTTGT	TTACATATTG	GAACAAGGCT	TCAACATCTG	AACCTATGGA	TTTTTTTACA 2000
2001	ATATCTGAAG	TCTGCCCTGCA	CCAGTTCCAG	TACATGGGGA	AGCCATACAT	AGCCAGGAAC	CAGGAGGGGT	TGGGACCCAT 2080
2081	AGTTCATGAT	CGAAAGTCTC	AGACATTGCC	TGTTTCCCGT	AACAGAAACAG	GAATGATGCA	TGCCAGATTG	CAGCAGCTGG 2160
2161	CGAGCCTGGA	TAACCTCTCT	ACTTTTAACC	ACAGCTATGG	CCACTCGGAC	GCAGATGTTT	TGCACCACTG	ATTACTTGAA 2240
2241	GCCAACATTG	CTACTGAGGT	TTGCCTGACA	GCTCTGGACA	CGCTTTCTCT	ATTTACATTG	GCGTTTAAGA	ACCAGCTCCT 2320
2321	GGCCGACCAT	GGACATAATC	CTCTCATGAA	AAAAGTTTTT	GATGTCTACC	TGTGTTTTCT	TCAAAAACAT	CAGTCTGAAA 2400
2401	CGGCTTTAAA	AAATGTCTTC	ACTGCCTTAA	GGTCCTTAAT	TTATAAGTTT	CCCTCAACAT	TCTATGAAGG	GAGAGCGGAC 2480
2481	ATGTGTGCGG	CTCTGTGTTA	CGAGATTCTC	AAGTGTCTGA	ACTCCAAGCT	GAGCTCCATC	AGGACGGAGG	CCTCCAGCT 2560
2561	GCTCTACTTC	CTGATGAGGA	ACAACCTTGA	TTACACTGGA	AAGAAGTCCT	TTGTCCGGAC	ACATTTGCAA	GTCATCATAT 2640
2641	CTGTACGCCA	GCTGATAGCA	GACGTTGTTG	GCATTGGGGA	AACCAGATTG	CAGCAGTCCC	TGTCATCAT	CAACAACCTG 2720
2721	GCCAACAGTG	ACCGGCTTAT	TAAGCACACC	AGCTTCTCCT	CTGATGTGAA	GGACTTAACC	AAAAGGATAC	GCACGGTGCT 2800
2801	AATGGCCACC	GCCCAGATGA	AGGAGCATGA	GAACGACCCA	GAGATGCTGG	TGGACCTCCA	GTACAGCCTG	GCCAAATCCT 2880
2881	ATGCCAGCAC	GCCCAGCTC	AGGAAGACGT	GGCTCGACAG	CATGGCCAGG	ATCCATGTCA	AAAATGGCGA	TCTCTCAGAG 2960
2961	CGCAATGAT	GCTATGTCCA	CGTAACAGCC	CTAGTGGCAG	AATATCTCAC	ACGGAAAGGC	GTGTTTAGAC	AAGGATGCAC 3040
3041	CGCCTTCAGG	GTCATTACCC	CAACATCGA	CGAGGAGGCC	TCCATGATGG	AAGACGTGGG	GATGCAGGAT	GTCCATTTCA 3120
3121	ACGAGGATGT	GCTGATGGAG	CTCCTTGAGC	AGTGCAGAGA	TGGACTCTGG	AAAGCCGAGC	GCTACGAGCT	CATCGCCGAC 3200
3201	ATCTACAAAC	TTATCATCCC	CATTTATGAG	AAGCGGAGGG	ATTTCTTTGA	AGATGAAGAT	GGAAAGGAGT	ATATTTACAA 3280
3281	GGAACCCAAA	CTCACACCGC	TGTCGGAAAT	TTCTCAGAGA	CTCCTTAAAC	TGTACTCGGA	TAAATTTGGT	TCTGAAAATG 3360
3361	TCAAAATGAT	ACAGGATTCT	GGCAAGGTCA	ACCCTAAGGA	TCTGGATTCT	AAGTATGCAT	ACATCCAGGT	GACTCACGTC 3440
3441	ATCCCCTTCT	TTGACGAAAA	AGAGTTGCAA	GAAAGGAAAA	CAGAGTTTGA	GAGATCCCAC	AACATCCGCC	GCTTCATGTT 3520
3521	TGAGATGCCA	TTTACGCAGA	CCGGGAAGAG	GCAGGGCGGG	GTGGAAGAGC	AGTGCAAACG	GCGCACCATC	CTGACAGCCA 3600
3601	TACACTGCTT	CCCTTATGTG	AAGAAGCGCA	TCCCTGTCT	GTACCAGCAC	CACACTGACC	TGAACCCCAT	CGAGGTGGCC 3680
3681	ATTGACGAGA	TGAGTAAGAA	GGTGGCGGAG	CTCCGGCAGC	TGTGCTCCTC	GGCCGAGGTG	GACATGATCA	AACTGCAGCT 3760
3761	CAAACTCCAG	GGCAGCGTGA	GTGTTCAAGT	CAATGCTGGC	CCACTAGCAT	ATGCGCGAGC	TTTCTTAGAT	GATACAAACA 3840
3841	CAAAGCGATA	TCCTGACAAT	AAAGTGAAGC	TGCTTAAGGA	AGTTTTCAGG	CAATTTGTGG	AAGCTTGCGG	TCAAGCCTTA 3920
3921	GCGGTAAACG	AACGTCTGAT	TAAAGAAGAC	CAGCTCGAGT	ATCAGGAAGA	AATGAAAGCC	AACTACAGGG	AAATGGCGAA 4000
4001	GGAGCTTTCT	GAAATCATGC	ATGAGCAGAT	CTGCCCCCTG	GAGGAGAAGA	CGAGCGTCTT	ACCGAATTCC	CTTCACATCT 4080
4081	TCAACGCCAT	CAGTGGGACT	CCAACAAGCA	CAATGGTTCA	CGGGATGACC	AGCTCGTCTT	CGGTGCTGTG	ATTACATCTC 4160
4161	ATGGCCCGTG	TGTGGGGACT	TGCTTTGTCA	TTTGCAAACT	CAGGATGCTT	TCCAAGGCCA	ATCACTGGGG	AGACCGAGCA 4240
4241	CAGGGAGGAC	CAAGGGGAAG	GGGAGAGAAA	GGAAATAAAG	AACAACGTTA	TTTCTTAACA	GACTTTCTAT	AGGAGTTGTA 4320
4321	AGAAGGTGCA	CATATTTTTT	TAAATCTCAC	TGGCAATATT	CAAAGTTTTT	ATTGTGTCTT	AACAAAGGTG	TGGTAGACAC 4400
4401	TCTTGAGCTG	GACTTAGATT	TTATTCTTCC	TTGCAGAGTA	GTGTTAGAAT	AGATGGCCTA	CAGAAAAAAA	AGGTTCTGGG 4480
4481	ATCTACATGG	CAGGGAGGGC	TGCACTGACA	TTGATGCCTG	GGGGACCTTT	TGCCTCGACT	CGTGCCGGAA	ATCTGATCGT 4560
4561	AATCAGGGTA	CAGAACTTAC	TAGTTTTGTC	TAGGAGTATG	TTGTATGACT	AGGATTTGTG	CTATTATCTC	ATTCAACAAC 4640
4641	ATAGAGCAAG	AATAGTGAGC	TAACTGAGCT	AGACACTCAA	TTAATCCGCT	ACTGGCTTCA	AGTCAGAACT	TTGTCATTAA 4720
4721	TCATCGACTC	CGGGACGGTC	ATATATGTAT	TACATTTCTA	CATTTTTAAT	ACTCACATGG	GCTTATGCAT	TAAGTTTAAT 4800
4801	TGTGATAAAT	TTGTGCTGGT	CCAGTATATG	CAATACACTT	TAATGGTTTA	TTCTTGTCAT	AAAAATGTGC	AATATGGAGA 4880
4881	TGTATACAAG	TCTTTACT						4898

FIG. 10A



	10	20	30	40	50	60	70	80
1	AATTGTAATA	CGACTCACTA	TAGGGCGAAT	TGGGTACCGG	GCCCCCCTC	GAGGTCGACG	GTATCGATAA	GCTTGATATC 80
81	GAATTTCGGCA	CGAGTTTTAC	ACCATCACCA	AAACCCAGAA	TTTTATGATG	AGATTAAAAT	AGAGTTGCCC	ACTCAGCTGC 160
161	ATGAAAAGCA	CCACCTGTTG	CTCACATTCT	TCCATGTCAG	CTGTGACAAC	TCAAGTAAAG	GAAGCACGAA	GAAGAGGGAT 240
241	GTCGTTGAAA	CCCAAGTTGG	CTACTCCTGG	CTTCCCCTCC	TGAAAGACGG	AAGGGTGGTG	ACAAGCGAGC	AGCACATCCC 320
321	GGTCTCGGCG	AACCTTCCTT	CGGGCTATCT	TGGCTACCAA	GAGCTTGGGA	TGGGCAGGCA	TTATGGTCCG	GAAATTAAAT 400
401	GGGTAGATGG	AGGCAAGCCA	CTGCTGAAAA	TTTCCACTCA	TCTGGTTTCT	ACAGGGATAC	TCAGGATCAG	CATTTACATA 480
481	ATTTTTTCCA	GTACTGTCAG	AAAACCGAAT	CTGGAGCCCA	AGCCTTAGGA	AACGAACCTG	TAAAGTACCT	TAAGAGTCTG 560
561	CATGCGATGG	AAGGCCACGT	GATGATCGCC	TTCTTGCCCA	CTATCCTAAA	CCAGCTGTTT	CGAGTCTCA	CCAGAGCCAC 640
641	ACAGGAAGAA	GTCGCGGTTA	ACGTGACTCG	GGTCATTATT	CATGTGGTTG	CCCAGTGCCA	TGAGGAAGGA	TTGGAGAGCC 720
721	ACTTGAGGTC	ATATGTTAAG	TACGCGTATA	AGGCTGAGCC	ATATGTTGCC	TCTGAATACA	AGACAGTGCA	TGAAGAAGTG 800
801	ACCAAATCCA	TGACCACGAT	TCTCAAGCCT	TCTGCCGATT	TCCTCACCAG	CAACAACTA	CTGAGGTACT	CATGGTTTTT 880
881	CTTTGATGTA	CTGATCAAAT	CTATGGCTCA	GCATTTGATA	GAGAACTCCA	AAGTTAAGTT	GCTGCGAAAC	CAGAGATTTT 960
961	CTGCATCCTA	TCATCATGCA	GCGGAAACCG	TTGTAAATAT	GCTGATGCCA	CACATCACTC	AGAACTTTGG	AGATAATCCA 1040
1041	GAGGCATCTA	AGAAGCGGAA	TCATAGCCTT	GCTGTCTTCA	TCAAGAGATG	TTTCACCTTC	ATGGACAGGG	GCTTTGTCTT 1120
1121	CAAGCAGATC	AACAATACA	TTAGCTGTTT	TGCTCCTGGA	GACCCAAAGA	CCCTCTTGA	ATACAAGTTT	GAATTTCTCC 1200
1201	GTGTAGTGTG	CAACCATGAA	CATTATATTC	CGTTGAACCT	ACCAATGCCA	TTTGGAAAAG	GCAGGATTCA	AAGATACCAA 1280
1281	GACCTCCAGC	TTGACTACTC	ATTAACAGAT	GAGTTCTGCA	GAAACCACTT	CTTGGTGGGA	CTGTTACTGA	GGGAGGTGGG 1360
1361	GACAGCCCTC	CAGGAGTTCC	GGGAGGTCCG	TCTGATCGCC	ATCAGTGTGC	TCAAGAACCT	GCTGATAAAG	CATTCTTTTG 1440
1441	ATGACAGATA	TGCTTCAAGG	AGCCATCAGG	CAAGGATAGC	CACCTCTAC	CTGCTCTGT	TTGGTCTGCT	GATTGAAAAC 1520
1521	GTCCAGCGGA	TCAATGTGAG	GGATGTGTCA	CCCTTCCCTG	TGAACGCGGG	CATGACCGTG	AAGGATGAAT	CCCTGGCTCT 1600
1601	ACCAGCTGTG	AATCCGCTGG	TGACGCCGCA	GAAGGGAAGC	ACCCTGGACA	ACAGCCTGCA	CAAGGACCTG	CTGGGCGCCA 1680
1681	TCTCCGSCAT	TGCTTCTCCA	TATACAACCT	CAACTCCAAA	CATCAACAGT	GTGAGAAATG	CTGATTGAG	AGGATCTCTC 1760
1761	ATAAGCACAG	ATTCGGGTAA	CAGCCTTCCA	GAAAGGAATA	GTGAGAAGAG	CAATTCCTTG	GATAAGCACC	AACAAAGTAG 1840
1841	CACATTGGGA	AATTCCGTGG	TTGCTGTGTA	TAAACTTGAC	CAGTCTGAGA	TTAAGAGCCT	ACTGATGTGT	TTCTCTTACA 1920
1921	TCTTAAAGAG	CATGTCTGAT	GATGCTTGT	TTACATATTG	GAACAAGGCT	TCAACATCTG	AACCTATGGA	TTTTTTTACA 2000
2001	ATATCTGAAG	TCTGCCTGCA	CCAGTTCAG	TACATGGGGA	AGCGATACAT	AGCCAGGAAC	CAGGAGGGGT	TGGGACCCAT 2080
2081	AGTTCATGAT	CGAAAGTCTC	AGACATTGCC	TGTTTCCCGT	AACAGAACAG	GAATGATGCA	TGCCAGATTG	CAGCAGCTGG 2160
2161	GCACCTGGGA	TAACTCTCTC	ACTTTTAACC	ACAGCTATGG	CCACTCGGAC	GCAGATGTTT	TGCACCAGTC	ATTACTTGAA 2240
2241	GCCAACATTG	CTACTGAGGT	TTGCCGTGACA	GCTCTGGACA	CGCTTTCTCT	ATTTACATTG	GCGTTTAAGA	ACCAGCTCCT 2320
2321	GGCCGACCAT	GGACATAATC	CTCTCATGAA	AAAAGTTTTT	GATGTCTACC	TGTGTTTTCT	TCAAAAACAT	CAGTCTGAAA 2400
2401	CGGCTTTAAA	AAATGTCTTC	ACTGCCTTAA	GGTCCTTAAT	TTATAAGTTT	CCCTCAACAT	TCTATGAAGG	GAGAGCGGAC 2480
2481	ATGTGTGCGG	CTCTGTGTTA	CGAGATTCTC	AAGTGTCTGA	ACTCCAAGCT	GAGCTCCATC	AGGACGGAGG	CCTCCCAGCT 2560
2561	GCTCTACTTC	CTGATGAGGA	ACAACCTTGA	TTACACTGGA	AAGAAGTCCT	TTGTCCGGAC	ACATTTGCAA	GTCTATCATAT 2640
2641	CTGTGAGCCA	GCTGATAGCA	GACGTTGTTG	GCATTGGGGA	AACCAGATTG	CAGCAGTCCC	TGTCCATCAT	CAACAACGTG 2720
2721	GCCAACAGTG	ACCGGCTTAT	TAAGCACACC	AGCTTCTCCT	CTGATGTGAA	GGACTTAACC	AAAAGGATAC	GCACGGTGCT 2800
2801	AATGGCCACC	GCCCAGATGA	AGGAGCATGA	GAACGACCCA	GAGATGCTGG	TGGACCTCCA	GTACAGCCTG	GCCAAATCCT 2880
2881	ATGCCAGCAC	GCCCGAGCTC	AGGAAGACGT	GGCTCGACAG	CATGGCCAGG	ATCCATGTCA	AAAATGGCGA	TCTCTCAGAG 2960
2961	GCAGCAATGT	GCTATGTCCA	CGTAACAGCC	CTAGTGGCAG	AATATCTCAC	ACGGAAGGCG	GTGTTTAGAC	AAGGATGCAC 3040
3041	CGCCTTCAGG	GTCATTACCC	CAAACATCGA	CGAGGAGGCC	TCCATGATGG	AAGACGTGGG	GATGCAGGAT	GTCCATTTCA 3120
3121	ACGAGGATGT	GCTGATGGAG	CTCCTTGAGC	AGTGCGCAGA	TGGACTCTGG	AAAGCCGAGC	GCTACGAGCT	CATCGCCGAC 3200
3201	ATCTACAAAC	TTATCATCCC	CATTTATGAG	AAGCGGAGGG	ATTTCTTTGA	AGATGAAGAT	GGAAAGGAGT	ATATTTTACA 3280
3281	GTAAACCCAA	CTCACACCGC	TGTGCGAAAT	TTCTCAGAGA	CTCCTTAAAC	TGTACTCGGA	TAAATTTGGT	TCTGAAAATG 3360
3361	TCAAAATGAT	ACAGGATTCT	GGCAAGGTCA	ACCCTAAGGA	TCTGGATTCT	AAGTATGCAT	ACATCCAGGT	GACTCACGTC 3440
3441	ATCCCCCTCT	TTGACGAAAA	AGAGTTGCAA	GAAAGGAAAA	CAGAGTTTGA	GAGATCCAC	AACATCCGCC	GCTTCATGTT 3520
3521	TGAGATGCCA	TTTACGCAGA	CCGGGAAGAG	GCAGGGCGGG	GTGGAAGAGC	AGTGCAAACG	GCGCACCATC	CTGACAGCCA 3600
3601	TACACTGCTT	CCCTTATGTG	AAGAAGCGCA	TCCCTGTCTAT	GTACCAGCAC	CACACTGACC	TGAACCCCAT	CGAGGTGGCC 3680
3681	ATTGACGAGA	TGAGTAAGAA	GGTGGCGGAG	CTCCGGCAGC	TGTGCTCCTC	GGCCGAGGTG	GACATGATCA	AACTGCAGCT 3760
3761	CAAACCTCCAG	GGCAGCGTGA	GTGTTCAAGT	CAATGCTGGC	CCACTAGCAT	ATGCGCGAGC	TTTCTTAGAT	GATACAAACA 3840
3841	CAAAGCGATA	TCCTGACAAT	AAAGTGAAGC	TGCTTAAGGA	AGTTTTTCAGG	CAATTTGTGG	AAGCTTGCGG	TCAAGCCTTA 3920
3921	GCGGTAAACG	AACGTCTGAT	TAAAGAAGAC	CAGCTCGAGT	ATCAGGAAGA	AATGAAAGCC	AACTACAGGG	AAATGGCGAA 4000
4001	GGAGCTTTCT	GAAATCATGC	ATGAGCAGAT	CTGCCCCCTG	GAGGAGAAGA	CGAGCGTCTT	ACCGAATTCC	CTTCACATCT 4080
4081	TCAACGCCAT	CAGTGGGACT	CCAACAAGCA	CAATGGTTCA	CGGGATGACC	AGCTCGTCTT	CGGTGCTGTG	ATTACATCTC 4160
4161	ATGGCCCGTG	TGTGGGGACT	TGCTTTGTCA	TTTGCAAACCT	CAGGATGCTT	TCCAAAGCCA	ATCACTGGGG	AGACCGAGCA 4240
4241	CAGGGAGGAC	CAAGGGGAAG	GGGAGAGAAA	GGAAATAAAG	AACAACGTTA	TTTCTTAACA	GACTTTCTAT	AGGAGTTGTA 4320
4321	AGAAGGTGCA	CATATTTTTT	TAAATCTCAC	TGGCAATATT	CAAAGTTTTT	ATTGTGTCTT	AACAAAGGTG	TGGTAGACAC 4400
4401	TCTTGAGCTG	GACTTAGATT	TTATTTCTCC	TTGCAGAGTA	GTGTTAGAAT	AGATGGCCTA	CAGAAAAAAA	AGGTTCTGGG 4480
4481	ATCTACATGG	CAGGGAGGGC	TGCACTGACA	TTGATGCCTG	GGGGACCTTT	TGCCTCGACT	CGTGCCGGAA	ATCTGATCGT 4560
4561	AATCAGGGTA	CAGAACTTAC	TAGTTTTGTC	TAGGAGTATG	TTGTATGACT	AGGATTTGTG	CTATTATCTC	ATTCAACAAC 4640
4641	ATAGAGCAAG	AATAGTGAGC	TAACTGAGCT	AGACACTCAA	TTAATCCGCT	ACTGGCTTCA	AGTCAGAACT	TTGTCAATTAA 4720
4721	TCATCGACTC	CGGGACGGTC	ATATATGTAT	TACATTTCTA	CATTTTTAAT	ACTCACATGG	GCTTATGCAT	TAAGTTTAAAT 4800
4801	TGTGATAAAT	TTGTGCTGGT	CCAGTATATG	CAATACACTT	TAATGGTTTA	TTCTGTCTAT	AAAAATGTGC	AATATGGAGA 4880
4881	TGTATACAAG	TCTTTACT						4898

FIG. 10B





	10	20	30	40	50	60	70	80	
1	AATTGTAATA	CGACTCACTA	TAGGGCGAAT	TGGGTACCGG	GCCCCCCTC	GAGGTGACG	GTATCGATAA	GCTTGATATC	80
1	GAATTCGGCA	CGAGTTTAC	ACCATCACCA	AAACCCAGAA	TTTTATGATG	AGATTAAAT	AGAGTTGCCC	ACTCAGCTGC	160
1	ATGAAAAGCA	CCACCTGTTG	CTCACATTCT	TCCATGTCAG	CTGTGACAA	TCAAGTAAAG	GAAGCACGAA	GAAGAGGGAT	240
11	GTCGTTGAAA	CCCAAGTTGG	CTACTCCTGG	CTTCCCCTCC	TGAAAGACGG	AAGGGTGGTG	ACAAGCGAGC	AGCACATCCC	320
21	GGTCTCGGCG	AACCTTCCTT	CGGGCTATCT	TGGCTACCAA	GAGCTTGGGA	TGGGCAGGCA	TTATGGTCCG	GAAATTAAAT	400
31	GGGTAGATGG	AGGCAAGCCA	CTGCTGAAAA	TTTCCACTCA	TCTGGTTTCT	ACAGGGATAC	TCAGGATCAG	CATTTACATA	480
41	ATTTTTTCCA	GTACTGTCAG	AAAACCGAAT	CTGGAGCCCA	AGCCTTAGGA	AACGAACCTG	TAAAGTACCT	TAAGAGTCTG	560
51	CATGCGATGG	AAGGCCACGT	GATGATCGCC	TTCTTGCCCA	CTATCCTAAA	CCAGCTGTTT	CGAGTCTCTA	CCAGAGCCAC	640
61	ACAGGAAGAA	GTCGCGGTTA	ACGTGACTCG	GGTCATTATT	CATGTGGTTG	CCCAGTGCCA	TGAGGAAGGA	TTGGAGAGCC	720
71	ACTTGAGGTC	ATATGTTAAG	TACGCGTATA	AGGCTGAGCC	ATATGTTGCC	TCTGAATACA	AGACAGTGCA	TGAAGAAGTG	800
81	ACCAAATCCA	TGACCACGAT	TCTCAAGCCT	TCTGCCGATT	TCCTCACCAG	CAACAACTA	CTGAGGTACT	CATGGTTTTT	880
91	CTTTGATGTA	CTGATCAAAT	CTATGGCTCA	GCATTTGATA	GAGAACTCCA	AAGTTAAGTT	GCTGCGAAAC	CAGAGATTTT	960
101	CTGCATCCTA	TCATCATGCA	GCGGAAACCG	TTGTAAATAT	GCTGATGCCA	CACATCACTC	AGAAGTTTGG	AGATAATCCA	1040
111	GAGGCATCTA	AGAACGCGAA	TCATAGCCTT	GCTGTCTTCA	TCAAGAGATG	TTTACCTTC	ATGGACAGGG	GCTTTGTCTT	1120
121	CAAGCAGATC	AACAACCTACA	TTAGCTGTTT	TGCTCCTGGA	GACCCAAAGA	CCCTCTTGA	ATACAAGTTT	GAATTTCTCC	1200
131	GTGTAGTGTG	CAACCATGAA	CATTATATTC	CGTTGAACCT	ACCAATGCCA	TTTGGAAAAG	GCAGGATTCA	AAGATACCAA	1280
141	GACCTCCAGC	TTGACTACTC	ATTAACAGAT	GAGTTCTGCA	GAAACCACTT	CTTGGTGGGA	CTGTTACTGA	GGGAGGTGGG	1360
151	GACAGCCCTC	CAGGAGTTCC	GGGAGGTCCG	TCTGATCGCC	ATCAGTGTGC	TCAAGAACCT	GCTGATAAAG	CATTCCTTTG	1440
161	ATGACAGATA	TGCTTCAAGG	AGCCATCAGG	CAAGGATAGC	CACCTCTAC	CTGCCCTGT	TTGGTCTGCT	GATTGAAAAC	1520
171	GTCCAGCGGA	TCAATGTGAG	GGATGTGTC	CCCTTCCCTG	TGAACGCGGG	CATGACCGTG	AAGGATGAAT	CCCTGGCTCT	1600
181	ACCAGCTGTG	AATCCGCTGG	TGACGCGGCA	GAAGGGAAGC	ACCTTGGACA	ACAGCTGCA	CAAGGACCTG	CTGGGCGCCA	1680
191	TCTCCGGCAT	TGCTTCTCCA	TATACAACCT	CAACTCCAAA	CATCAACAGT	GTGAGAAATG	CTGATTCGAG	AGGATCTCTC	1760
201	ATAAGCACAG	ATTCCGGTAA	CAGCCTTCCA	GAAAGGAATA	GTGAGAAGAG	CAATTCCCTG	GATAAGCACC	AACAAAGTAG	1840
211	CACATTGGGA	AATTCCGTGG	TTGCTGTGTA	TAAACTTGAC	CAGTCTGAGA	TTAAGAGCCT	ACTGATGTGT	TTCTCTTACA	1920
221	TCTTAAAGAG	CATGTCTGAT	GATGCTTTGT	TTACATATTG	GAACAAGGCT	TCAACATCTG	AACTTATGGA	TTTTTTTACA	2000
231	ATATCTGAAG	TCTGCCTGCA	CCAGTTCAG	TACATGGGGA	AGCGATACAT	AGCCAGGAAC	CAGGAGGGGT	TGGGACCCAT	2080
241	AGTTTATGAT	CGAAAGTCTC	AGACATGACC	TGTTTCCCGT	AACAGAACAG	GAATGATGCA	TGCCAGATTG	CAGCAGCTGG	2160
251	GGAGCCTGGA	TAACTCTCTC	ACTTTTAAAC	ACAGCTATGG	CCACTCGGAC	GCAGATGTTT	TGCACCAGTC	ATTACTTGAA	2240
261	GCCAACATTG	CTACTGAGGT	TTGCCTGACA	GCTCTGGACA	CGCTTTCTCT	ATTTACATTG	GCGTTTAAAG	ACCAGCTCCT	2320
271	GGCCGACCAT	GGACATAATC	CTCTCATGAA	AAAAGTTTTT	GATGTCTACC	TGTGTTTTCT	TCAAAAACAT	CAGTCTGAAA	2400
281	CGGCTTTAAA	AAATGTCTTC	ACTGCCTTAA	GGTCTTTAAT	TTATAAGTTT	CCCTCAACAT	TCTATGAAGG	GAGAGCGGAC	2480
291	AGTGTGTGCG	CTCTGTGTTA	CGAGATTCTC	AAGTGTGTTA	ACTCCAAGCT	GAGCTCCATC	AGGACGGAGG	CCTCCAGCTT	2560
301	GCTCTACTTC	CTGATGAGGA	ACAACCTTGA	TTACACTGGA	AAGAAGTCTT	TGTCCGGAC	ACATTTGCAA	GTCATCATAT	2640
311	CTGTACGCCA	GCTGATAGCA	GACGTTGTTG	GCATTGGGGA	AACCAGATTC	CAGCAGTCCC	TGTCCATCAT	CAACAACGTG	2720
321	GCCAACAGTG	ACCGGCTTAT	TAAGCACACC	AGCTTCTCCT	CTGATGTGAA	GGACTTAACC	AAAAGGATAC	GCACGGTGCT	2800
331	AATGGCCACC	GCCCAGATGA	AGGAGCATGA	GAACGACCCA	GAGATGCTGG	TGGACCTCCA	GTACAGCCTG	GCCAAATCCT	2880
341	ATGCCAGCAC	GCCCAGAGCTC	AGGAAGACGT	GGCTCGACAG	CATGGCCAGG	ATCCATGTCA	AAAATGGCGA	TCTCTCAGAG	2960
351	GCAGCAATGT	GCTATGTCCA	CGTAACAGCC	CTAGTGGCAG	AATATCTCAC	ACGGAAGGCG	GTTGTTAGAC	AAGGATGCAC	3040
361	CGCCTTCAGG	GTCATTACCC	CAAACATCGA	CGAGGAGGCC	TCCATGATGG	AAGACGTGGG	GATGCAGGAT	GTCCATTTCA	3120
371	ACGAGGATGT	GCTGATGGAG	CTCCTTGAGC	AGTGCGCAGA	TGGACTCTGG	AAAGCCGAGC	GCTACGAGCT	CATCGCCGAC	3200
381	ATCTACAAAA	TTATCATCCC	CATTATGAG	AAGCGGAGGG	ATTTCTTTGA	AGATGAAGAT	GGAAAGGAGT	ATATTTACAA	3280
391	GGAAACCAAA	CTCACACCGC	TGTGCGAAAT	TTCTCAGAGA	CTTCTTAAAC	TGTACTCGGA	TAAATTTGGT	TCTGAAAATG	3360
401	TCAAAATGAT	ACAGGATTCT	GGCAAGGTCA	ACCCTAAGGA	TCTGGATTCT	AAGTATGCAT	ACATCCAGGT	GACTCACGTC	3440
411	ATCCCTTTCT	TTGACGAAAA	AGAGTTGCAA	GAAAGGAAAA	CAGAGTTTGA	GAGATCCAC	AACATCCGCC	GCTTCATGTT	3520
421	TGAGATGCCA	TTTACGCGAGA	CCGGGAAGAG	GCAGGGCGGG	GTGGAAGAGC	AGTGCAACG	GCGCACCATC	CTGACAGCCA	3600
431	TACACTGCTT	CCCTTATGTG	AAGAAGCGCA	TCCCTGTCTAT	GTACCAGCAC	CACACTGACC	TGAACCCCAT	CGAGGTGGCC	3680
441	ATTGACGAGA	TGAGTAAGAA	GGTGGCGGAG	CTCCGGCAGC	TGTGCTCCTC	GGCCGAGGTG	GACATGATCA	AACTGCAGCT	3760
451	CAAACCTCCAG	GGCAGCGTGA	GTGTTCAAGT	CAATGCTGGC	CCACTAGCAT	ATGCCGCGAGC	TTTCTTAGAT	GATACAAACA	3840
461	CAAAGCGATA	TCCTGACAAT	AAAGTGAAGC	TGCTTAAGGA	AGTTTTTCAGG	CAATTTGTGG	AAGCTTGCGG	TCAAGCCTTA	3920
471	GCGGTAAACG	AACGTCTGAT	TAAAGAAGAC	CAGCTCGAGT	ATCAGGAAGA	AATGAAAGCC	AACTACAGGG	AAATGGCGAA	4000
481	GGAGCTTTCT	GAAATCATGC	ATGAGCAGAT	CTGCCCCCTG	GAGGAGAAGA	CGAGCGTCTT	ACCGAATTCC	CTTCACATCT	4080
491	TCAACGCCAT	CAGTGGGACT	CCAACAAGCA	CAATGGTTCA	CGGGATGACC	AGCTCGTCTT	CGTCTGTTGT	ATTACATCTC	4160
501	ATGGCCCGTG	TGTGGGGACT	TGCTTTGTCA	TTTGCAAACCT	CAGGATGCTT	TCCAAAGCCA	ATCACTGGGG	AGACCGAGCA	4240
511	CAGGGAGGAC	CAAGGGGAAG	GGGAGAGAAA	GGAAATAAAG	AACAACGTTA	TTTCTTAACA	GACTTTCTAT	AGGAGTTGTA	4320
521	AGAAGGTGCA	CATATTTTTT	TAAATCTCAC	TGGCAATATT	CAAAGTTTTT	ATTGTGTCTT	AACAAAGGTG	TGTTAGACAC	4400
531	TCTTGAGCTG	GACTTAGATT	TTATTCTTCC	TTGCAGAGTA	GTGTTAGAAT	AGATGGCCTA	CAGAAAAAAA	AGGTTCTGGG	4480
541	ATCTACATGG	CAGGGAGGGC	TGCACTGACA	TTGATGCCTG	GGGGACCTTT	TGCTCGACT	CGTGCCGGAA	ATCTGATCGT	4560
551	AATCAGGGTA	CAGAACTTAC	TAGTTTGTGC	TAGGAGTATG	TTGTATGACT	AGGATTTGTG	CTATTATCTC	ATTCAACAAC	4640
561	ATAGAGCAAG	AATAGTGAGC	TAACTGAGCT	AGACACTCAA	TTAATCCGCT	ACTGGCTTCA	AGTCAGAACT	TTGTCATTAA	4720
571	TCATCGACTC	CGGGACGGTC	ATATATGTAT	TACATTTCTA	CATTTTTTAAT	ACTCACATGG	GCTTATGCAT	TAAGTTTAAT	4800
581	TGTGATAAAT	TTGTGCTGGT	CCAGTATATG	CAATACACTT	TAATGGTTTA	TTCTGTCTAT	AAAAATGTGC	AATATGGAGA	4880
591	TGTATACAAG	TCCTTACT							4898

FIG. 10C

	10	20	30	40	50	60	70	80
MEGHVMI AFL	PTILNQLFRV	LTRATQEEVA	VNVTRVIIHV	VAQCHEEGLE	SHLRSYVKYA	YKAEPYVASE	YKTVHEELTK	80
SMTTILKPSA	DFLTSTNKLRL	YSWFFFDVLI	KSMAGHLIEN	SRVKLLRNQR	FPASYHHAAE	TVVMMLMPHI	TQKPGDNPEA	160
SKVANHSLAV	FIKRCFTFMD	RGFVFKQINN	YISCFAPGDP	KTLFEYKFEF	LRVVCNHEHY	IPLNLPMPFG	KGRIQRYQDL	240
QLDYSLTDEF	CRNHFLVGLL	LREVGTALQE	FREVRLIAIS	VLKONLLIKHS	FDDRYASRSH	QARIATLYLP	LFGLLIENVQ	320
RINVRDVSPF	PVNAGMTVKD	ESLALPAVNP	LVTPQKGSTL	DNSLHKDLLG	AISGIASPYT	TSTPNINSVR	NADSRGSLIS	400
TDSGNSLPER	NSEKSNLSDK	HQOSSTLGNS	VVRCDKLDQS	EIKSLIMCFL	YILKMSDDA	LFTYWNKAST	SELMOFFTIS	480
EVCLHQFQYM	GKRYIARNQE	GLGPIVHDRK	SQTLFVSRNR	TGMMHARLQQ	LGSIDNSLTF	NHSYGHSDAD	VLHQSLLEAN	560
IATEVCLTAL	DTLSLFTLAF	KNQLLADHGH	NPLMKKVFDV	YLCFLQKHQS	ETALKNVFTA	LRSLIYKFPS	TFYEGRADMC	640
AALCYEILKC	CNSKLSSIRT	EASQLLYFLM	RNNFDYTGGK	SFVRTHLQVI	ISVSQLIADV	VGIGETRPOQ	SLSIINNCAN	720
SDRLIKHTSF	SSDVKDLTKR	IRTVLMATAQ	MKEHENDPEM	LVDLQYSLAK	SYASTPELRK	TWLDSEMARH	VKNGDLSEAA	800
MCYVHV TALV	AEYLTRKGVF	RQGCTAFRVI	TPNIDEEASM	MEDVGMQDVH	FNEDVLMELL	EQCADGLWKA	ERYELIADIY	880
KLIPIYEKR	RDFFEDGDK	EYIYKEPKLT	PLSEISQRL	KLYSDKFGSE	NVKMIQDSGK	VNPKDLDSKY	AYIQVTHVIP	960
FFDEKELQER	KTEFERSHNI	RRFMFEMPFT	QTGKRQGGVE	EQCKRTILT	AIHCFPYVKK	RIPVMYQHHT	DLNPIEVAID	1040
EMSKVAELR	QLCSSAEVDM	IKLQKLQGS	VSVQVNAGPL	AYARAFIDDT	NTKRYPDNKV	KLLKEVFRQF	VEACQALAV	1120
NERLIKEDQL	EYQCEMKANY	REMAKELSEI	MHEQICPLEE	KTSVLPNSLH	IFNAISGTPT	STMVHGMTSS	SSVV	1194

00687837 401300

FIG. 10C (cont.)

	10	20	30	40	50	60	70	80
1	AATTGTAATA	CGACTCACTA	TAGGGCGAAT	TGGGTACCGG	GCCCCCCCCTC	GAGGTGACG	GTATCGATAA	GCTTGATATC 80
81	GAATTCGGCA	CGAGTTTAC	ACCATCACCA	AAACCCAGAA	TTTTATGATG	AGATTAAAT	AGAGTTGCC	ACTCAGCTGC 160
61	ATGAAAAGCA	CCACCTGTTG	CTCACATTCT	TCCATGTCAG	CTGTGACAAC	TCAAGTAAAG	GAAGCACGAA	GAAGAGGGAT 240
41	GTCGTTGAAA	CCCAAGTTGG	CTACTCCTGG	CTTCCCCTCC	TGAAAGACGG	AAGGGTGGTG	ACAAGCGAGC	AGCACATCCC 320
21	GGTCTCGGCG	AACCTTCCTT	CGGGCTATCT	TGGCTACCAA	GAGCTTGGGA	TGGGCAGGCA	TTATGGTCCG	GAAATTAAAT 400
01	GGGTAGATGG	AGGCAAGCCA	CTGCTGAAAA	TTTCCACTCA	TCTGGTTTCT	ACAGGGATAC	TCAGGATCAG	CATTTACATA 480
81	ATTTTTTCCA	GTACTGTCAG	AAAACCGAAT	CTGGAGCCCA	AGCCTTAGGA	AACGAACCTG	TAAAGTACCT	TAAGAGTCTG 560
61	CATGCGATGG	AAGGCCACGT	GATGATCGCC	TTCTTGCCCA	CTATCCTAAA	CCAGCTGTTT	CGAGTCCTCA	CCAGAGCCAC 640
541	ACAGGAAGAA	GTGCGGTTA	ACGTGACTCG	GGTCATTATT	CATGTGGTTG	CCCAGTGCCA	TGAGGAAGGA	TTGGAGAGCC 720
721	ACTTGAGGTC	ATATGTTAAG	TACGCGTATA	AGGCTGAGCC	ATATGTTGCC	TCTGAATACA	AGACAGTGCA	TGAAGAACTG 800
801	ACCAAATCCA	TGACCACGAT	TCTCAAGCCT	TCTGCCGATT	TCCTCACCAG	CAACAACTA	CTGAGGTACT	CATGGTTTTT 880
881	CTTTGATGTA	CTGATCAAAT	CTATGGCTCA	GCATTTGATA	GAGAACTCCA	AAGTTAAGTT	GCTGCGAAAC	CAGAGATTTC 960
961	CTGCATCCTA	TCATCATGCA	GCGGAAACCG	TTGTAATAT	GCTGATGCCA	CACATCACTC	AGAAGTTTGG	AGATAATCCA 1040
041	GAGGCATCTA	AGAACGCGAA	TCATAGCCTT	GCTGTCTTCA	TCAAGAGATG	TTTACCTTC	ATGGACAGGG	GCTTTGTCTT 1120
121	CAAGCAGATC	AACAACCTACA	TTAGCTGTTT	TGCTCCTGGA	GACCCAAAGA	CCCTCTTTGA	ATACAAGTTT	GAATTTCTCC 1200
201	GTGTAGTGTG	CAACCATGAA	CATTATATTC	CGTTGAACCT	ACCAATGCCA	TTTGGAAAAG	GCAGGATTCA	AAGATACCAA 1280
281	GACCTCCAGC	TTGACTACTC	ATTAACAGAT	GAGTTCTGCA	GAAACCACTT	CTTGGTGGGA	CTGTTACTGA	GGGAGGTGGG 1360
1361	GACAGCCCTC	CAGGAGTTCC	GGGAGGTCCG	TCTGATCGCC	ATCAGTGTGC	TCAAGAACCT	GCTGATAAAG	CATTCTTTTG 1440
1441	ATGACAGATA	TGCTTCAAGG	AGCCATCAGG	CAAGGATAGC	CACCTCTAC	CTGCCCTCTG	TTGGTCTGCT	GATTGAAAAC 1520
1521	GTCCAGCGGA	TCAATGTGAG	GGATGTGTCA	CCCTTCCCTG	TGAACGCGGG	CATGACCGTG	AAGGATGAAT	CCCTGGCTCT 1600
1601	ACCAGCTGTG	AATCCGCTGG	TGACGCGCA	GAAGGGAAGC	ACCCTGGACA	ACAGCCTGCA	CAAGGACCTG	CTGGGCGCCA 1680
1681	TCTCCGGCAT	TGCTTCTCCA	TATACAACCT	CAACTCCAAA	CATCAACAGT	GTGAGAAATG	CTGATTGAG	AGGATCTCTC 1760
1761	ATAAGCACAG	ATTCCGGTAA	CAGCCTTCCA	GAAAGGAATA	GTGAGAAGAG	CAATTCCCTG	GATAAGCACC	AACAAAGTAG 1840
1841	CACATTGGGA	AATTCOGTGG	TTCGCTGTGA	TAAACTTGAC	CAGTCTGAGA	TTAAGAGCCT	ACTGATGTGT	TTCTCTTACA 1920
1921	TCTTAAAGAG	CATGTCTGAT	GATGCTTTGT	TTACATATTG	GAACAAGGCT	TCAACATCTG	AACCTATGGA	TTTTTTTACA 2000
2001	ATATCTGAAG	TCTGCCTGCA	CCAGTCCAG	TACATGGGGA	AGCGATACAT	AGCCAGGAAC	CAGGAGGGGT	TGGGACCCAT 2080
2081	AGTTCATGAT	CGAAAGTCTC	AGACATTGCC	TGTTTCCCGT	AACAGAACAG	GAATGATGCA	TGCCAGATTG	CAGCAGCTGG 2160
2161	GCAGCCTGGA	TAACCTCTCT	ACTTTTAACC	ACAGCTATGG	CCACTCGGAC	GCAGATGTTT	TGCACCACTG	ATTACTTGAA 2240
2241	GCCAACATTG	CTACTGAGGT	TTGCCGACA	GCTCTGGACA	CGCTTCTCT	ATTATCATTG	GCGTTTAAGA	ACCAGCTCCT 2320
2321	GGCCGACCAT	GGACATAATC	CTCTCATGAA	AAAAGTTTTT	GATGTTCTAC	TGTGTTTTCT	TCAAAAACAT	CAGTCTGAAA 2400
2401	GGGCTTTAAA	AAATGCTTTC	ACTGCCTTAA	GGTCTTAAT	TTATAAGTTT	CCCTCAACAT	TCTATGAAGG	GAGAGCGGAC 2480
2481	ATGTGTGCGG	CTCTGTGTTA	CGAGATTCTC	AAGTGCTGTA	ACTCCAAGCT	GAGCTCCATC	AGGACGGAGG	CCTCCCAGCT 2560
2561	GCTCTACTTC	CTGATGAGGA	ACAACCTTGA	TTACACTGGA	AAGAAGTCTT	TTGTCCGGAC	ACATTTGCAA	GTCATCATAT 2640
2641	CTGTACGCCA	GCTGATAGCA	GACGTTGTG	GCATTGGGGA	AACCAGATTC	CAGCAGTCCC	TGTCCATCAT	CAACAACCTG 2720
2721	GCCAACAGTG	ACCGGCTTAT	TAAGCACACC	AGCTTCTCCT	CTGATGTGAA	GGACTTAACC	AAAAGGATAC	GCACGGTGCT 2800
2801	AATGGCCACC	GCCCAGATGA	AGGAGCATGA	GAAAGACCCA	GAGATGCTGG	TGGACCTCCA	GTACAGCCTG	GCCAAATCCT 2880
2881	ATGCCAGCAC	GCCCAGGCTC	AGGAAGACGT	GGCTCGACAG	CATGGCCAGG	ATCCATGTCA	AAAATGGCGA	TCTCTCAGAG 2960
2961	GCAGCAATGT	GCTATGTCCA	CGTAACAGCC	CTAGTGGCAG	AATATCTCAC	ACGGAAGGCG	GTGTTTAGAC	AAGGATGCAC 3040
3041	CGCCTTCAGG	GTCATTACCC	CAAACATCGA	CGAGGAGGCC	TCCATGATGG	AAGACGTGGG	GATGCAGGAT	GTCCATTTC 3120
3121	ACGAGGATGT	GCTATGGAG	CTCCTTGAGC	AGTGCCGAGA	TGGACTCTGG	AAAGCCGAGC	GCTACGAGCT	CATCGCCGAC 3200
3201	ATCTACAAAC	TTATCATCCC	CATTATGAG	AAGCGGAGGG	ATTCTTTTGA	AGATGAAGAT	GGAAAGGAGT	ATATTACAA 3280
3281	GGAAACCAAA	CTCACACCGC	TGTCGGAAAT	TTCTCAGAGA	CTCCTTAAAC	TGTACTCGGA	TAAATTGGGT	TCTGAAAATG 3360
3361	TCAAAATGAT	ACAGGATTCT	GGCAAGGTCA	ACCCTAAGGA	TCTGGATTCT	AAGTATGCAT	ACATCCAGGT	GACTCAOGTC 3440
3441	ATCCCTTTCT	TTGACGAAAA	AGAGTTGCAA	GAAAGGAAAA	CAGAGTTTGA	GAGATCCAC	AACATCCGCC	GCTTCATGTT 3520
3521	TGAGATGCCA	TTTACGCGA	CCGGGAAGAG	GCAGGGCGGG	GTGGAAGAGC	AGTGCAAACG	GCGCACCATC	CTGACAGCCA 3600
3601	TACACTGCTT	CCCTTATGTG	AAGAAGCGCA	TCCCTGTCTAT	GTACCAGCAC	CACACTGACC	TGAACCCCAT	CGAGGTGGCC 3680
3681	ATTGACGAGA	TGAGTAAGAA	GGTGGCGGAG	CTCCGGCAGC	TGTGCTCCTC	GGCCGAGGTG	GACATGATCA	AACTGCAGCT 3760
3761	CAAACCTCCAG	GGCAGCGTGA	GTGTTCAAGT	CAATGCTGGC	CCACTAGCAT	ATGCGCGAGC	TTTCTTAGAT	GATACAAACA 3840
3841	CAAAGCGATA	TCTTGACAAT	AAAGTGAAGC	TGCTTAAGGA	AGTTTTTCAAG	CAATTTGTGG	AAGCTTGCGG	TCAAGCCTTA 3920
3921	GCGGTAAACG	AACGTCTGAT	TAAAGAAGAC	CAGCTCGAGT	ATCAGGAAGA	AATGAAAGCC	AACTACAGGG	AAATGGCGAA 4000
4001	GGAGCTTTCT	GAAATCATGC	ATGAGCAGAT	CTGCCCCCTG	GAGGAGAAGA	CGAGCGTCTT	ACCGAATTCC	CTTCACATCT 4080
4081	TCAACGCCAT	CAGTGGGACT	CCAACAAGCA	CAATGGTTCA	CGGGATGACC	AGCTCGTCTT	CGGTGCTGTG	ATTACATCTC 4160
4161	ATGGCCCGTG	TGTGGGGACT	TGCTTTGTCA	TTTGCAAACT	CAGGATGCTT	TCCAAGCCA	ATCACTGGGG	AGACCGAGCA 4240
4241	CAGGGAGGAC	CAAGGGGAAG	GGGAGAGAAA	GGAAATAAAG	AACAACGTTA	TTTCTTAACA	GACTTTCTAT	AGGAGTTGTA 4320
4321	AGAAGGTGCA	CATATTTTTT	TAAATCTCAC	TGGCAATATT	CAAAGTTTTT	ATTGTGCTT	AACAAAGGTG	TGGTAGACAC 4400
4401	TCTTGAGCTG	GACTTAGATT	TTATTTCTCC	TTGCAGAGTA	GTGTTAGAAT	AGATGGCCTA	CAGAAAAAAA	AGGTTCTGGG 4480
4481	ATCTACATGG	CAGGGAGGGC	TGCACTGACA	TTGATGCCCTG	GGGGACCTTT	TGCTCGACT	CGTGCCGGAA	ATCTGATCGT 4560
4561	AATCAGGGTA	CAGAACTTAC	TAGTTTTGTC	TAGGAGTATG	TTGTATGACT	AGGATTTGTG	CTATTATCTC	ATTCAACAAC 4640
4641	ATAGAGCAAG	AATAGTGAGC	TAACTGAGCT	AGACACTCAA	TTAATCCGCT	ACTGGCTTCA	AGTCAGAACT	TTGTCATTAA 4720
4721	TCATCGACTC	CGGGACGGTC	ATATATGTAT	TACATTTCTA	CATTTTTAAT	ACTCACATGG	GCTTATGCAT	TAAGTTTAAT 4800
4801	TGTGATAAAT	TTGTGCTGGT	CCAGTATATG	CAATACACTT	TAATGGTTTA	TTCTTGTCAT	AAAAATGTGC	AATATGGAGA 4880
4881	TGTATACAAG	TCTTTACT						4898

FIG. 10D

	10	20	30	40	50	60	70	80
MEGHVMI AFL	PTILNQLFRV	LTRATQEEVA	VNVTRVTHV	VAQCHEEGLE	SHLSYVKYA	YKAEFYVASE	YKTVHEELTK	80
SMTTILKPSA	DFLTSNKLLR	YSWFFFDVLI	KSMAQH LIEN	SKVKLLRNQR	FPASYHAAE	TVVNM LMPHI	TQKFGDNPEA	160
SKNANHSLAV	FIKRCFTFMD	RGFVFKQINN	YISCFAPGDP	KTLFEYKFEF	LRVVCNHEHY	IPLNLPMPFG	KGRIQRYQDL	240
QLDYSLTDEF	CRNHFLVGLL	LREVG TALQE	FREVLIAIS	VLK NLLIKHS	FDDRYASRSH	QARIATLYLP	LFGLLIENVQ	320
RINVRDVSPF	PVNAGMTVKD	ESLALPAVNP	LVTPQKGSTL	DNSLHKDLIG	AISGIASPYT	TSTPNINSVR	NADSRGSLIS	400
1 TDSGNSLPER	NSEKSNSLDK	HQQSSTLGNS	VVRCDKLDQS	EIKSLMCFL	YILKSMSDDA	LFTYWNKAST	SELMDFFTIS	480
1 EVCLHQFOYM	GKRYIARNQE	GLGPIVHDRK	SQTLFVSRNR	TGMMHARLQQ	LGSLDNSLTF	NHSYGHSDAD	VLHQSLLEAN	560
1 IATEVCLTAL	DTLSLFTLAF	KNQLLADHGH	NPLMKKVFDV	YLCFLOKHQS	ETALKNVFTA	LRSLIYKFPS	TFYEGRADMC	640
1 AALCYEILKC	CNSKLSSIRT	EASQLLYFLM	RNNFDYTGKK	SFVRTHLQVI	ISVSQLIADV	VGIGETRFOQ	SLSIINN CAN	720
1 SDRLIKHTSF	SSDVKDLTKR	IRTVLMATAQ	MKEHENDPEM	LVDLOYS LAK	SYASTPELRK	TWLD SMARIH	VKNGDLSEAA	800
1 MCYVHV TALV	AEYLTRKGVF	RQGCTAFRVI	TPNIDEEASM	MEDVGMQDVH	FNEDVIMELL	EQCADGLWKA	ERYELIADIY	880
1 KLIPIY EKR	RDFPEDEDGK	EYIYKEPKLT	PLSEISQRLI	KLYSDKFGSE	NVKMIQDSGK	VNPKDLDSKY	AYIQVTHVIP	960
51 FFDEKELQER	KTEFERSHNI	RRFMFEMPFT	QTGKRQGGVE	EQCKRTILT	AIHCFPYVKK	RIPVMYQHHT	DLNPIEVAID	1040
41 EMSKVAELR	QLCSSAEVDM	IKLQLKLQGS	VSVQVNAGPL	AYARAFLDDT	NTRYPDNKV	KLLKEVFRQF	VEACQALAV	1120
21 NERLIKEDQL	EYQEEMKANY	REMAKELSEI	MHEQICPLEE	KTSVLPNSLH	IFNAISGTPT	STMVHGMTSS	SSVV	1194

FIG. 10D (cont.)



	10	20	30	40	50	60	70	80
AATTGTAATA	CGACTCACTA	TAGGGCGAAT	TGGGTACCGG	GCCCCCCTC	GAGGTGACG	GTATCGATAA	GCTTGATATC	80
GAATTCGGCA	CGAGTTTAC	ACCATCACCA	AAACCCAGAA	TTTTATGATG	AGATTAAAT	AGAGTTGCC	ACTCAGCTGC	160
ATGAAAAGCA	CCACCTGTTG	CTCACATTCT	TCCATGTCAG	CTGTGACAAC	TCAAGTAAAG	GAAGCACGAA	GAAGAGGGAT	240
GTCGTTGAAA	CCCAAGTTGG	CTACTCCTGG	CTTCCCCCTC	TGAAAGACGG	AAGGGTGGTG	ACAAGCGAGC	AGCACATCCC	320
GGTCTCGGCG	AACCTTCCTT	CGGGCTATCT	TGGCTACCAA	GAGCTTGGGA	TGGGCAGGCA	TTATGGTCCG	GAAATTAAAT	400
GGGTAGATGG	AGGCAAGCCA	CTGCTGAAAA	TTTCCACTCA	TCTGGTTTCT	ACAGGGATAC	TCAGGATCAG	CATTTACATA	480
ATTTTTTCCA	GTACTGTCAG	AAAACCGAAT	CTGGAGCCCA	AGCCTTAGGA	AACGAACTTG	TAAAGTACCT	TAAGAGTCTG	560
CATGCGATGG	AAGGCCACGT	GATGATCGCC	TTCTTGCCCA	CTATCCTAAA	CCAGCTGTTT	CGAGTCCTCA	CCAGAGCCAC	640
ACAGGAAGAA	GTCGCGGTTA	ACGTGACTCG	GGTCATTATT	CATGTGGTTG	CCCAGTGCCA	TGAGGAAGGA	TTGGAGAGCC	720
ACTTGAGGTC	ATATGTTAAG	TACGCGTATA	AGGCTGAGCC	ATATGTTGCC	TCTGAATACA	AGACAGTGCA	TGAAGAACTG	800
ACCAAATCCA	TGACCACGAT	TCTCAAGCCT	TCTGCCGATT	TCCTCACCAG	CAACAACTA	CTGAGGTACT	CATGGTTTTT	880
CTTTGATGTA	CTGATCAAAT	CTATGGCTCA	GCATTTGATA	GAGAACTCCA	AAGTTAAGTT	GCTGCGAAAC	CAGAGATTTC	960
CTGCATCCTA	TCATCATGCA	GCGGAAACCG	TTGTAATAT	GCTGATGCCA	CACATCACTC	AGAAGTTTGG	AGATAATCCA	1040
GAGGCATCTA	AGAACGCGAA	TCATAGCCTT	GCTGTCTTCA	TCAAGAGATG	TTTCACCTTC	ATGGACAGGG	GCTTTGTCTT	1120
CAAGCAGATC	AACAACCTACA	TTAGCTGTTT	TGCTCCTGGA	GACCCAAAGA	CCCTCTTTGA	ATACAAGTTT	GAATTTCTCC	1200
GTGTAGTGTG	CAACCATGAA	CATTATATTC	CGTTGAACTT	ACCAATGCCA	TTTGGAAAAG	GCAGGATTCA	AAGATACCAA	1280
GACCTCCAGC	TTGACTACTC	ATTAACAGAT	GAGTTCTGCA	GAAACCACTT	CTTGGTGGGA	CTGTTACTGA	GGGAGGTGGG	1360
GACAGCCCTC	CAGGAGTTCC	GGGAGGTCCG	TCTGATCGCC	ATCAGTGTGC	TCAAGAACCT	GCTGATAAAG	CATTCTTTTG	1440
ATGACAGATA	TGCTTCAAGG	AGCCATCAGG	CAAGGATAGC	CACCTCTAC	CTGCCCTGT	TTGGTCTGCT	GATTGAAAAC	1520
GTCCAGCGGA	TCAATGTGAG	GGATGTGTCA	CCCTTCCCTG	TGAACGCGGG	CATGACCGTG	AAGGATGAAT	CCCTGGCTCT	1600
ACCAGCTGTG	AATCCGCTGG	TGACGCGCGA	GAAGGGAAGC	ACCCTGGACA	ACAGCCTGCA	CAAGGACCTG	CTGGGCGCCA	1680
TCTCCGGCAT	TGCTTCTCCA	TATACAACTT	CAACTCCAAA	CATCAACAGT	GTGAGAAATG	CTGATTGAG	AGGATCTCTC	1760
ATAAGCACAG	ATTCCGGTAA	CAGCCTTCCA	GAAAGGAATA	GTGAGAAGAG	CAATTCCCTG	GATAAGCACC	AACAAAGTAG	1840
CACATTGGGA	AATTCGCTGG	TTCCGCTGTA	TAAACTTGAC	CAGTCTGAGA	TTAAGAGCCT	ACTGATGTGT	TTCTCTTACA	1920
TCTTAAAGAG	CATGTCTGAT	GATGCTTTGT	TTACATATTG	GAACAAGGCT	TCAACATCTG	AACTTATGGA	TTTTTTTACA	2000
ATATCTGAAG	TCTGCCTGCA	CCAGTTCCAG	TACATGGGGA	AGCGATACAT	AGCCAGGAAC	CAGGAGGGGT	TGGGACCCAT	2080
AGTTTCATGAT	CGAAAGTCTC	AGACATTGCC	TGTTTCCCGT	AACAGAACAG	GAATGATGCA	TGCCAGATTG	CAGCAGCTGG	2160
GCAGCTGGA	TAACTCTCTC	ACTTTTAACC	ACAGCTATGG	CCACTCGGAC	GCAGATGTTT	TGCACCAGTC	ATTACTTGAA	2240
GCCACATTG	CTACTGAGGT	TTGCCTGACA	GCTCTGGACA	CGCTTCTCT	ATTACATTG	GCGTTTAAGA	ACCAGCTCCT	2320
GGCGAACCAT	GGACATAATC	CTCTCATGAA	AAAAGTTTTT	GATGTCTACC	TGTGTTTTCT	TCAAAAACAT	CAGTCTGAAA	2400
CGGCTTTAAA	AAATGCTTTC	ACTGCCTTAA	GGTCCCTAAT	TTATAAGTTT	CCCTCAACAT	TCTATGAAGG	GAGAGCGGAC	2480
ATGTGTGCGG	CTCTGTGTTA	CGAGATTCTC	AAGTGCTGTA	ACTCCAAGCT	GAGCTCCATC	AGGACGGAGG	CCTCCAGCT	2560
GCTCTACTTC	CTGATGAGGA	ACAACCTTGA	TTACACTGGA	AAGAAGTCTT	TGTCCGGAC	ACATTGCA	GTCATCATAT	2640
CTGTGAGCCA	GCTGATAGCA	GACGTTGTTG	GCATTGGGGA	AACCAGATTC	CAGCAGTCCC	TGTCCATCAT	CAACAACGTG	2720
GCCACAGTG	ACCGGCTTAT	TAAGCACACC	AGCTTCTCCT	CTGATGTGAA	GGACTTAACC	AAAAGGATAC	GCACGGTGCT	2800
AATGGCCACC	GCCCAGATGA	AGGAGCATGA	GAACGACCCA	GAGATGCTGG	TGGACCTCCA	GTACAGCCTG	GCCAAATCCT	2880
ATGECAGCAC	GCCCAGCTC	AGGAAGACGT	GGCTCGACAG	CATGGCCAGG	ATCCATGTCA	AAAATGGCGA	TCTCTCAGAG	2960
GCAGCAATGT	GCTATGTCCA	CGTAACAGCT	CTAGTGGCAG	AATATCTCAC	ACGGAAGGC	GTGTTTAGAC	AAGGATGCAC	3040
CGCCTTCAGG	GTCATTACCC	CAAACATCGA	CGAGGAGGCC	TCCATGATGG	AAGACGTGGG	GATGCAGGAT	GTCCATTTC	3120
ACGAGGATGT	GCTGATGGAG	CTCCTTGAGC	AGTGCGCAGA	TGGACTCTGG	AAAGCCGAGC	GCTACGAGCT	CATCGCCGAC	3200
ATETACAAAC	TTATCATCCC	CATTATGAG	AAGCGGAGGG	ATTTCTTTGA	AGATGAAGAT	GGAAAGGAGT	ATATTTACAA	3280
GGAAACCCAA	CTCACACCGC	TGTCCGAAAT	TTCTCAGAGA	CTCCTTAAAC	TGTACTCGGA	TAAATTTGGT	TCTGAAAATG	3360
TCAAAATGAT	ACAGGATTCT	GGCAAGGTCA	ACCCTAAGGA	TCTGGATTCT	AAGTATGCAT	ACATCCAGGT	GACTCACGTC	3440
ATCCCCCTCT	TTGACGAAAA	AGAGTTGCAA	GAAAGGAAAA	CAGAGTTTGA	GAGATCCAC	AACATCCGCC	GCTTCATGTT	3520
TGAGATGCCA	TTTACGCGAG	CCGGGAAGAG	GCAGGGCGGG	GTGGAAGAGC	AGTGCAAACG	GCGCACCATC	CTGACAGCCA	3600
TACTACTGCTT	CCCTTATGTG	AAGAAGCGCA	TCCCTGTCTAT	GTACCAGCAC	CACACTGACC	TGAACCCCAT	CGAGGTGGCC	3680
ATTGACGAGA	TGAGTAAGAA	GGTGGCGGAG	CTCCGGCAGC	TGTGCTCCTC	GGCCGAGGTG	GACATGATCA	AACTGCAGCT	3760
CAACTCCAG	GGCAGCGTGA	GTGTTCAAGT	CAATGCTGGC	CCACTAGCAT	ATGCGCGAGC	TTTCTTAGAT	GATACAAACA	3840
CAAAGCGATA	TCTGACAAAT	AAAGTGAAGC	TGCTTAAGGA	AGTTTTCAGG	CAATTTGTGG	AAGCTTGCGG	TCAAGCCTTA	3920
GCGGTAAACG	AACGTCTGAT	TAAAGAAGAC	CAGCTCGAGT	ATCAGGAAGA	AATGAAAGCC	AACTACAGGG	AAATGGCGAA	4000
GGAGCTTTCT	GAAATCATGC	ATGAGCAGAT	CTGCCCCCTG	GAGGAGAAGA	CGAGCGTCTT	ACCGAATTCC	CTTCACATCT	4080
TCAACGCCAT	CAGTGGGACT	CCAACAAGCA	CAATGGTTCA	CGGGATGACC	AGCTCGTCTT	CGGTGCTGTG	ATTACATCTC	4160
ATGGCCCGTG	TGTGGGGACT	TGCTTTGTCA	TTTGCAAACT	CAGGATGCTT	TCCAAAGCCA	ATCACTGGGG	AGACCGAGCA	4240
CAGGGAGGAC	CAAGGGGAAG	GGGAGAGAAA	GGAAATAAAG	AACAACGTTA	TTTCTTAAAC	GACTTTCTAT	AGGAGTTGTA	4320
AGAAGGTGCA	CATATTTTTT	TAAATCTCAC	TGGCAATATT	CAAAGTTTTT	ATTGTGTCTT	AACAAAGGTG	TGGTAGACAC	4400
TCTTGAGCTG	GACTTAGATT	TTATTCTTCC	TTGCAGAGTA	GTGTTAGAAT	AGATGGCCTA	CAGAAAAAAA	AGGTTCTGGG	4480
ATCTACATGG	CAGGGAGGGC	TGCACTGACA	TTGATGCCTG	GGGGACCTTT	TGCCTCGACT	CGTGCCGGAA	ATCTGATCGT	4560
AATCAGGGTA	CAGAACTTAC	TAGTTTTGTC	TAGGAGTATG	TTGTATGACT	AGGATTTGTG	CTATTATCTC	ATTCAACAAC	4640
ATAGAGCAAG	AATAGTGAGC	TAAGTGAGCT	AGACACTCAA	TTAATCCGCT	ACTGGCTTCA	AGTCAGAACT	TTGTCAATTAA	4720
TCATCGACTC	CGGGACGGTC	ATATATGTAT	TACATTTCTA	CATTTTTAAT	ACTCACATGG	GCTTATGCAT	TAAGTTTAAAT	4800
TGTGATAAAT	TTGTGCTGGT	CCAGTATATG	CAATACACTT	TAATGGTTTA	TTCTTGTCAT	AAAAATGTGC	AATATGGAGA	4880
TGTATACAAG	TCITTTACT							4898

FIG. 10E



10	20	30	40	50	60	70	80
HVMIAFL	PTILNQLFRV	LTRATQEEVA	VNVTRVIIHV	VAQCHEEGLE	SHLRSYVKYA	YKAEPYVASE	YKTVHEELTK 80
TILKPSA	DFLTENKLLR	YSWFFFDVLI	KSMAQH LIEN	SKVKLLRNQR	FPASYHHAAE	TVVNMIMPHI	TQKPGDNPEA 160
IANHSLAV	FIKRCFTFMD	RGFVFKQINN	YISCFAPGDP	KTLFEYKFEF	LRVVCNHEHY	IFLNLPMPPG	KGRIQRYQDL 240
YSLTDEF	CRNHFLVGLL	LREVGTAQOE	FREVRLIAIS	VLNLLIKHS	FDDRYASRSH	QARIATLYLP	LPGLLIENVQ 320
VRDVSPF	PVNAGMTVID	ESLALPAVNP	LVTPOKGSTL	DNSLHKDLIG	AISGIASPYT	TSTPNINSVR	NADSRGSLIS 400
SGNSLPER	NSEKSNSLDK	HQOSSTLGNS	VVRCDKLDQS	EIKSLIMCFL	YILKSMDDA	LFTYWNKAST	SEIMDFFTIS 480
CLHQFOYM	GKRYIARNQE	GLGPIVHDRK	SQTLFVSRNR	TGMHARLOQ	LGSLDNLSTF	NHSYGHSDAD	VLHQSLLEAN 560
TEVCLTAL	DTLSLFTLAF	KNQLLADHGH	NPLMKKVFDV	YLCFLQKHQS	ETALKQVFTA	LRSLIYKFPS	TFYEGRADMC 640
LCYEILKC	CNSKLSSIRT	EASQLLYFLM	RNNFDYTGGK	SFVRTHLOVI	ISVSQLLADV	VGIGETRFOQ	SLSIDNNCAN 720
RLIKHTSF	SSDVKDLTKR	IRTVLMATAQ	MKEHENDPEM	LVDLOYSIAK	SYASTPELRK	TWLDSEARIH	VKNGDLSEAA 800
YVHTALV	AEYLTRKGVF	ROGCTAFRVI	TPNIDEEASM	MEDVGMQDVH	FNEDVIMELL	EQCADGLWKA	ERYELLADIY 880
AIPIYEKR	RDFFEDEDGK	EYIYKEPKLT	PLSEISORLL	KLYSDKFGSE	NVKMIQDSGK	VNPKDLDSKY	AYIQVTHVIP 960
DEKELQER	KTEFERSHNI	RRFMFEMPFT	QTGKRGQGVF	EQCKRTILT	AIHCFFYVKK	RIPVMYQHHT	DLNPIEVAID 1040
MSKKVAELR	QLCSSAEVDM	IKLQKLOGS	VSVQVWAGPL	AYARAFDDT	NTKRYPDNKV	KLLKEVFRQF	VEACQALAV 1120
ERLIKEDQL	EYQEEWKANY	REMAKELSEI	MHEQICPLEE	KTSVLPNSLH	IFKASGTPT	STMVHGMTSS	SSVV 1194

FIG. 1DE (cont.)

	10	20	30	40	50	60	70	80
AATTGTAATA	CGACTCACTA	TAGGGCGAAT	TGGGTACCGG	GCCCCCCTC	GAGGTGACG	GTATCGATAA	GCTTGATATC	80
GAATTCGGCA	CGAGTTTAC	ACCATCACCA	AAACCCAGAA	TTTTATGATG	AGATTAAAT	AGAGTTGCC	ACTCAGCTGC	160
ATGAAAAGCA	CCACCTGTTG	CTCACATTCT	TCCATGTCAG	CTGTGACAAC	TCAAGTAAAG	GAAGCACGAA	GAAGAGGGAT	240
GTCGTTGAAA	CCCAAGTTGG	CTACTCCTGG	CTTCCCTCC	TGAAAGACGG	AAGGGTGGTG	ACAAGCGAGC	AGCACATCCC	320
GGTCTCGGCG	AACCTTCCTT	CGGGCTATCT	TGGCTACCAA	GAGCTTGGGA	TGGGCAGGCA	TTATGGTCCG	GAAATTAAAT	400
GGGTAGATGG	AGGCAAGCCA	CTGCTGAAAA	TTTCCACTCA	TCTGGTTTCT	ACAGGGATAC	TCAGGATCAG	CATTTACATA	480
ATTTTTTCCA	GTACTGTCAG	AAAACCGAAT	CTGGAGCCCA	AGCCTTAGGA	AACGAACTTG	TAAAGTACCT	TAAGAGTCTG	560
CATGCGATGG	AAGGCCACGT	GATGATCGCC	TTCTTGCCCA	CTATCCTAAA	CCAGCTGTTT	CGAGTCTCTA	CCAGAGCCAC	640
ACAGGAAGAA	GTCGCGGTTA	ACGTGACTCG	GGTCATTATT	CATGTGGTTG	CCCAGTGCCA	TGAGGAAGGA	TTGGAGAGCC	720
ACTTGAGGTC	ATATGTTAAG	TACGCGTATA	AGGCTGAGCC	ATATGTTGCC	TCTGAATACA	AGACAGTGCA	TGAAGAACTG	800
ACCAAATCCA	TGACCACGAT	TCTCAAGCCT	TCTGCGGATT	TCCTCACCAG	CAACAACTA	CTGAGGTACT	CATGGTTTTT	880
CTTTGATGTA	CTGATCAAAT	CTATGGCTCA	GCATTTGATA	GAGAACTCCA	AAGTTAAGTT	GCTGCGAAAC	CAGAGATTTC	960
CTGCATCCTA	TCATCATGCA	GCGGAAACCG	TTGTAATAT	GCTGATGCCA	CACATCACTC	AGAAGTTTGG	AGATAATCCA	1040
GAGGCATCTA	AGAACGCGAA	TCATAGCCTT	GCTGTCTTCA	TCAAGAGATG	TTTCACCTTC	ATGGACAGGG	GCTTTGTCTT	1120
CAAGCAGATC	AACAACCTACA	TTAGCTGTTT	TGCTCCTGGA	GACCCAAAGA	CCCTCTTGA	ATACAAGTTT	GAATTTCTCC	1200
GTGTAGTGTG	CAACCATGAA	CATTATATTC	CGTTGAACCT	ACCAATGCCA	TTTGGAAAAG	GCAGGATTCA	AAGATACCAA	1280
GACCTCCAGC	TTGACTACTC	ATTAACAGAT	GAGTTCTGCA	GAAACCACTT	CTTGGTGGGA	CTGTTACTGA	GGGAGGTGGG	1360
GACAGCCCTC	CAGGAGTTCC	GGGAGGTCCG	TCTGATCGCC	ATCAGTGTGC	TCAAGAACCT	GCTGATAAAG	CATTCCTTTG	1440
ATGACAGATA	TGCTTCAAGG	AGCCATCAGG	CAAGGATAGC	CACCTCTAC	CTGCTCTGT	TTGGTCTGCT	GATTGAAAAC	1520
GTCAGCGGA	TCAATGTGAG	GGATGTGTCA	CCCTTCCCTG	TGAACGCGGG	CATGACCGTG	AAGGATGAAT	CCCTGGCTCT	1600
ACCAGCTGTG	AATCCGCTGG	TGACGCCGCA	GAAGGGAAGC	ACCCTGGACA	ACAGCCTGCA	CAAGGACCTG	CTGGGCGCCA	1680
TCTCCGGCAT	TGCTTCTCCA	TATACAACCT	CAACTCCAAA	CATCAACAGT	GTGAGAAATG	CTGATTCGAG	AGGATCTCTC	1760
ATAAGCACAG	ATTCCGGTAA	CAGCCTTCCA	GAAAGGAATA	GTGAGAAGAG	CAATTCCCTG	GATAAGCACC	AACAAAGTAG	1840
CACATTGGGA	AATTCGGTGG	TTGCTGTGTA	TAAACTTGAC	CAGTCTGAGA	TTAAGAGCCT	ACTGATGTGT	TTCTCTTACA	1920
TCITAAAGAG	CATGTCTGAT	GATGCTTTGT	TTACATATTG	GAACAAGGCT	TCAACATCTG	AACTTATGGA	TTTTTTTACA	2000
ATATCTGAAG	TCTGCCTGCA	CCAGTTCCAG	TACATGGGGA	AGCGATACAT	AGCCAGGAAC	CAGGAGGGGT	TGGGACCCAT	2080
AGTTCATGAT	CGAAAGTCTC	AGACATTGCC	TGTTTCCCGT	AACAGAACAG	GAATGATGCA	TGCCAGATTG	CAGCAGCTGG	2160
GCAGCTGGGA	TAACTCTCTC	ACTTTTAAAC	ACAGCTATGG	CCACTCGGAC	GCAGATGTTT	TGCACCAGTC	ATTACTTGAA	2240
GCCACCATTG	CTACTGAGGT	TTGCTTGACA	GCTCTGGACA	CGCTTCTCT	ATTTACATTG	GCGTTTAAAG	ACCAGCTCCT	2320
GGCGGACCAT	GGACATAATC	CTCTCATGAA	AAAAGTTTTT	GATGTCTACC	TGTGTTTTCT	TCAAAAACAT	CAGTCTGAAA	2400
CGGCTTTAAA	AAATGTCTTC	ACTGCCTTAA	GGTCTTAAT	TTATAAGTTT	CCCTCAACAT	TCTATGAAGG	GAGAGCGGAC	2480
ATGTGTGCGG	CTCTGTGTTA	CGAGATTCTC	AAGTGCTGTA	ACTCCAAGCT	GAGCTCCATC	AGGACGGAGG	CCTCCAGCT	2560
GCTGTACTTC	CTGATGAGGA	ACAACCTTGA	TTACACTGGA	AAGAAGTCTT	TTGTCCGGAC	ACATTTGCAA	GTCATCATAT	2640
CTGTGAGCCA	GCTGATAGCA	GACGTGTGTT	GCATTGGGGA	AACCAGATT	CAGCAGTCCC	TGTCCATCAT	CAACAACGTG	2720
GCCAACAGTG	ACCGGCTTAT	TAAGCACACC	AGCTTCTCCT	CTGATGTGAA	GGACTTAACC	AAAAGGATAC	GCACGGTGCT	2800
AATGGCCACC	GCCCAGATGA	AGGAGCATGA	GAACGACCCA	GAGATGCTGG	TGGACCTCCA	GTACAGCCTG	GCCAAATCCT	2880
ATGCCAGCAC	GCCCGAGCTC	AGGAAGACGT	GGCTCGACAG	CATGGCCAGG	ATCCATGTCA	AAAATGGCGA	TCTCTCAGAG	2960
GCAGCAATGT	GCTATGTCCA	CGTAACAGCC	CTAGTGCCAG	AATATCTCAC	ACGGAAGGCC	GTGTTTAGAC	AAGGATGCAC	3040
CGCCTTCAGG	GTCATTACCC	CAAACATCGA	CGAGGAGGCC	TCCATGATGG	AAGACGTGGG	GATGCAGGAT	GTCCATTTC	3120
ACGAGGATGT	GCTGATGGAG	CTCCTTGAGC	AGTGCGCAGA	TGGACTCTGG	AAAGCCGAGC	GCTACGAGCT	CATCGCCGAC	3200
ATGTAACAAAC	TTATCATCCC	CATTTATGAG	AAGCGGAGGG	ATTTCTTTGA	AGATGAAGAT	GGAAAGGAGT	ATATTTACAA	3280
GGAAACCCAAA	CTCACACCGC	TGTCCGAAAT	TTCTCAGAGA	CTCCTTAAAC	TGTACTCGGA	TAAATTTGGT	TCTGAAAATG	3360
TCMAATGAT	ACAGGATTCT	GGCAAGGTCA	ACCCTAAGGA	TCTGGATTCT	AAGTATGCAT	ACATCCAGGT	GACTCACGTC	3440
ATGECCTTCT	TTGACGAAAA	AGAGTTGCAA	GAAAGGAAAA	CAGAGTTTGA	GAGATCCAC	AACATCCGCC	GCTTCATGTT	3520
TGAGATGCCA	TTTACGCGA	CCGGGAAGAG	GCAAGGCGGG	GTGGAAGAGC	AGTGCAACG	GCGACCCATC	CTGACAGCCA	3600
TACACTGCTT	CCCTTATGTG	AAGAAGCGCA	TCCCTGTCTAT	GTACCAGCAC	CACACTGACC	TGAACCCCAT	CGAGGTGGCC	3680
ATTTGACGAGA	TGAGTAAGAA	GGTGGCGGAG	CTCCGCGCAGC	TGTGCTCCTC	GGCCGAGGTG	GACATGATCA	AACTGCAGCT	3760
CAAACTCCAG	GGCAGCGTGA	GTGTTCAAGT	CAATGCTGGC	CCACTAGCAT	ATGCGCGAGC	TTTCTTAGAT	GATACAAACA	3840
CAAAGCGATA	TCCTGACAAT	AAAGTGAAGC	TGCTTAAGGA	AGTTTTTCAGG	CAATTTGTGG	AAGCTTGCGG	TCAAGCCTTA	3920
GCGGTAAACG	AACGTCTGAT	TAAAGAAGAC	CAGCTCGAGT	ATCAGGAAGA	AATGAAAGCC	AACTACAGGG	AAATGGCGAA	4000
GGAGCTTTCT	GAAATCATGC	ATGAGCAGAT	CTGCCCCCTG	GAGGAGAAGA	CGAGCGTCTT	ACCGAATTCC	CTTCACATCT	4080
TCAACGCCAT	CAGTGGGACT	CCAACAAGCA	CAATGGTTCA	CGGGATGACC	AGCTCGTCTT	CGGTGCTGTG	ATTACATCTC	4160
ATGGCCCGTG	TGTGGGGACT	TGCTTTGTCA	TTTGCAAAC	CAGGATGCTT	TCCAAAGCCA	ATCACTGGGG	AGACCGAGCA	4240
CAGGGAGGAC	CAAGGGGAAG	GGGAGAGAAA	GGAAATAAAG	AACAACGTTA	TTTCTTAAAC	GACTTTCTAT	AGGAGTTGTA	4320
AGAAGGTGCA	CATATTTTTT	TAAATCTCAC	TGGCAATATT	CAAAGTTTTC	ATTGTGTCTT	AACAAAGGTG	TGGTAGACAC	4400
TCTTGAGCTG	GACTTAGATT	TTATTCTTCC	TTGCAGAGTA	GTGTTAGAA	AGATGGCCTA	CAGAAAAAAA	AGGTTCTGGG	4480
ATCTACATGG	CAGGGAGGGC	TGCACTGACA	TTGATGCCTG	GGGGACCTTT	TGCCTCGACT	CGTGCCGGAA	ATCTGATCGT	4560
AATCAGGGTA	CAGAACTTAC	TAGTTTTGTC	TAGGAGTATG	TTGTATGACT	AGGATTTGTG	CTATTATCTC	ATTCAACAAC	4640
ATAGAGCAAG	AATAGTGAGC	TAACTGAGCT	AGACACTCAA	TTAATCCGCT	ACTGGCTTCA	AGTCAGAACT	TTGTCAATTAA	4720
TCATCGACTC	CGGGACGGTC	ATATATGTAT	TACATTTCTA	CATTTTAAAT	ACTCACATGG	GCTTATGCAT	TAAGTTTAAAT	4800
TGTGATAAAT	TTGTGCTGGT	CCAGTATATG	CAATACACTT	TAATGGTTTA	TTCTTGTCAT	AAAAATGTGC	AATATGGAGA	4880
TGTATACAAG	TCTTTACT							4898



	10	20	30	40	50	60	70	80
1	AATTGTAATA	CGACTCACTA	TAGGGCGAAT	TGGGTACCGG	GCCCCCCTC	GAGGTCGACG	GTATCGATAA	GCTTGATATC 80
81	GAATTCGGCA	CGAGTTTAC	ACCATCACCA	AAACCCAGAA	TTTTATGATG	AGATTAAAT	AGAGTTGCC	ACTCAGCTGC 160
161	ATGAAAAGCA	CCACCTGTTG	CTCACATTCT	TCCATGTCAG	CTGTGACAAC	TCAAGTAAAG	GAAGCACGAA	GAAGAGGGAT 240
241	GTCGTTGAAA	CCCAAGTTGG	CTACTCCTGG	CTTCCCCTCC	TGAAAGACGG	AAGGGTGGTG	ACAAGCGAGC	AGCACATCCC 320
321	GGTCTCGGCG	AACCTTCCTT	CGGGCTATCT	TGGCTACCAA	GAGCTTGGGA	TGGGCAGGCA	TTATGGTCCG	GAAATTAAAT 400
401	GGGTAGATGG	AGGCAAGCCA	CTGCTGAAAA	TTTCCACTCA	TCTGGTTTCT	ACAGGGATAC	TCAGGATCAG	CATTTACATA 480
481	ATTTTTTCCA	GTACTGTCAG	AAAACCGAAT	CTGGAGCCCA	AGCCTTAGGA	AACGAACCTG	TAAAGTACCT	TAAGAGTCTG 560
561	CATGCGATGG	AAGGCCACGT	GATGATCGCC	TTCTTGCCCA	CTATCCTAAA	CCAGCTGTTT	CGAGTCCTCA	CCAGAGCCAC 640
641	ACAGGAAGAA	GTCGCGGTTA	ACGTGACTCG	GGTCATTATT	CATGTGGTTG	CCCAGTGCCA	TGAGGAAGGA	TTGGAGAGCC 720
721	ACTTGAGGTG	ATATGTTAAG	TACGCGTATA	AGGCTGAGCC	ATATGTTGCC	TCTGAATACA	AGACAGTGCA	TGAAGAACTG 800
801	ACCAAATCCA	TGACCACGAT	TCTCAAGCCT	TCTGCCGATT	TCCTCACCAG	CAACAACTA	CTGAGGTACT	CATGGTTTTT 880
881	CTTTGATGTA	CTGATCAAAT	CTATGGCTCA	GCATTTGATA	GAGAACTCCA	AAGTTAAGTT	GCTGCGAAAC	CAGAGATTTC 960
961	CTGCATCCTA	TCATCATGCA	GCGGAAACCG	TTGTAAATAT	GCTGATGCCA	CACATCACTC	AGAAGTTTGG	AGATAATCCA 1040
1041	GAGGCATCTA	AGAACGCGAA	TCATAGCCTT	GCTGTCTTCA	TCAAGAGATG	TTTCACTTTC	ATGGACAGGG	GCTTTGTCTT 1120
1121	CAAGCAGATC	AACAACCTACA	TTAGCTGTTT	TGCTCCTGGA	GACCCAAAGA	CCCTCTTTGA	ATACAAGTTT	GAATTTCTCC 1200
1201	GTGTAGTGTG	CAACCATGAA	CATTATATTC	CGTTGAACTT	ACCAATGCCA	TTTGAAAAG	GCAGGATTCA	AAGATACCAA 1280
1281	GACCTCCAGC	TTGACTACTC	ATTAACAGAT	GAGTTCTGCA	GAAACCACTT	CTTGGTGGGA	CTGTTACTGA	GGGAGGTGGG 1360
1361	GACAGCCCTC	CAGGAGTTCC	GGGAGGTCCG	TCTGATCGCC	ATCAGTGTGC	TCAAGAACCT	GCTGATAAAG	CATTCTTTTG 1440
1441	ATGACAGATA	TGCTTCAAGG	AGCCATCAGG	CAAGGATAGC	CACCTCTAC	CTGCCCTGTG	TTGGTCTGCT	GATTGAAAAC 1520
1521	GTCCAGCGGA	TCAATGTGAG	GGATGTGTCA	CCCTTCCCTG	TGAACGCGGG	CATGACCCTG	AAGGATGAAT	CCCTGGCTCT 1600
1601	ACCAGCTGTG	AATCCGCTGG	TGACGCGCA	GAAGGGGAAGC	ACCTTGGACA	ACAGCCTGCA	CAAGGACCTG	CTGGGCGCCA 1680
1681	TCTCCGGCAT	TGCTTCTCCA	TATACAACCT	CAACTCCAAA	CATCAACAGT	GTGAGAAATG	CTGATTGAG	AGGATCTCTC 1760
1761	ATAAGCACAG	ATTCCGGTAA	CAGCCTTCCA	GAAAGGAATA	GTGAGAAGAG	CAATTCCTTG	GATAAGCACC	AACAAAGTAG 1840
1841	CACATTGGGA	AATTCCGTGG	TTGCTGTGA	TAAACTTGAC	CAGTCTGAGA	TTAAGAGCCT	ACTGATGTGT	TTCTCTTACA 1920
1921	TCTTAAAGAG	CATGTCTGAT	GATGCTTTGT	TTACATATTG	GAACAAGGCT	TCAACATCTG	AACCTTATGA	TTTTTTTACA 2000
2001	ATATCTGAAG	TCTGCCTGCA	CCAGTTCCAG	TACATGGGGA	AGCGATACAT	AGCCAGGAAC	CAGGAGGGGT	TGGGACCCAT 2080
2081	AGTTTCATGAT	CGAAAGTCTC	AGACATTGCC	TGTTTCCCGT	AACAGAACAG	GAATGATGCA	TGCCAGATTG	CAGCAGCTGG 2160
2161	GCAGCCTGGA	TAACTCTCTC	ACTTTTAACT	ACAGCTATGG	CCACTCGGAC	GCAGATGTTT	TGCACCAGTC	ATTACTTGAA 2240
2241	GCCAACATTG	CTACTGAGGT	TTGCCTGACA	GCTCTGGACA	CGCTTTCTCT	ATTACATTG	GCGTTTAAAG	ACCAGCTCCT 2320
2321	GGCCGACCAT	GGACATAATC	CTCTCATGAA	AAAAGTTTTT	GATGTCTACC	TGTGTTTTCT	TCAAAAACAT	CAGTCTGAAA 2400
2401	CGGCTTTAAA	AAATGTCTTC	ACTGCCITAA	GGTCTTAAAT	TTATAAGTTT	CCCTCAACAT	TCTATGAAGG	GAGAGCGGAC 2480
2481	ATGTGTGCGG	CTCTGTGTTA	CGAGATTCTC	AAGTGCTGTA	ACTCCAAGCT	GAGCTCCATC	AGGACGGAGG	CCTCCCAGCT 2560
2561	GCTCTACTTC	CTGATGAGGA	ACAACCTTGA	TTACTACTGGA	AAGAAGTCTT	TTGTCCGGAC	ACATTTGCAA	GTCATCATAT 2640
2641	CTGTACAGCA	GCTGATAGCA	GACGTTGTTG	GCATTGGGGA	AACCAGATTC	CAGCAGTCCC	TGTCCATCAT	CAACAACGTG 2720
2721	GCCAACAGTG	ACCGGCTTAT	TAAGCACACC	AGCTTCTCCT	CTGATGTGAA	GGACTTAAAC	AAAAGGATAC	GCACGGTGCT 2800
2801	AATGGCCACC	GCCCAGATGA	AGGAGCATGA	GAACGACCCA	GAGATGCTGG	TGGACCTCCA	GTACAGCCTG	GCCAAATCCT 2880
2881	ATGCCAGCAC	GCCCGAGCTC	AGGAAGACGT	GGCTCGACAG	CATGGCCAGG	ATCCATGTCA	AAAATGGCGA	TCTCTCAGAG 2960
2961	GCAGCAATGT	GCTATGTCCA	CGTAACAGCC	CTAGTGGCAG	AATATCTCAC	ACGGAAGGCG	GTGTTTAGAC	AAGGATGCAC 3040
3041	CGCCTTCAGG	GTCATTACCC	CAAACATCGA	CGAGGAGGCC	TCCATGATGG	AAGACGTGGG	GATGCAGGAT	GTCCATTTC A 3120
3121	ACGAGGATGT	GCTGATGGAG	CTCCTTGAGC	AGTGCGCAGA	TGGACTCTGG	AAAGCCGAGC	GCTACGAGCT	CATCGCCGAC 3200
3201	ATCTACAAAC	TTATCATCCC	CATTTATGAG	AAGCGGAGGG	ATTTCTTTGA	AGATGAAGAT	GGAAAGGAGT	ATATTTACAA 3280
3281	GGAACCCAAA	CTCACACCGC	TGTCGGAAAT	TTCTCAGAGA	CTCCTTAAAC	TGTACTCGGA	TAAATTTGGT	TCTGAAAATG 3360
3361	TCAAAATGAT	ACAGGATTCT	GGCAAGGTCA	ACCCTAAGGA	TCTGGATTCT	AAGTATGCAT	ACATCCAGGT	GACTCACGTC 3440
3441	ATCCCCTTCT	TTGACGAAAA	AGAGTTGCAA	GAAAGGAAAA	CAGAGTTTGA	GAGATCCCAC	AACATCCGCC	GCTTCATGTT 3520
3521	TGAGATGCCA	TTTACGCAGA	CCGGGAAGAG	GCAGGGCGGG	GTGGAAGAGC	AGTGCAAACG	GCGCACCATC	CTGACAGCCA 3600
3601	TACACTGCTT	CCCTTATGTG	AAGAAGCGCA	TCCCTGTCTAT	GTACCAGCAC	CACACTGACC	TGAACCCCAT	CGAGGTGGCC 3680
3681	ATTGACGAGA	TGAGTAAGAA	GGTGGCGGAG	CTCCGGCAGC	TGTGCTCCTC	GGCCGAGGTG	GACATGATCA	AACTGCAGCT 3760
3761	CAAACCTCCAG	GGCAGCGTGA	GTGTTCAAGT	CAATGCTGGC	CCACTAGCAT	ATGCGCGAGC	TTTCTTAGAT	GATACAAACA 3840
3841	CAAAGCGATA	TCCTGACAAT	AAAGTGAAGC	TGCTTAAGGA	AGTTTTTCAGG	CAATTTGTGG	AAGCTTGCGG	TCAAGCCTTA 3920
3921	GCGGTAAACG	AACGTCTGAT	TAAAGAAGAC	CAGCTCGAGT	ATCAGGAAGA	AATGAAAGCC	AACTACAGGG	AAATGGCGAA 4000
4001	GGAGCTTTCT	GAAATCATGC	ATGAGCAGAT	CTGCCCCCTG	GAGGAGAAGA	CGAGCGTCTT	ACCGAATTCC	CTTCACATCT 4080
4081	TCAACGCCAT	CAGTGGGACT	CCAACAAGCA	CAATGGTTCA	CGGGATGACC	AGCTCGTCTT	CGGTGCTGTG	ATTACATCTC 4160
4161	ATGGCCCGTG	TGTGGGGACT	TGCTTTGTCA	TTTGCAAACT	CAGGATGCTT	TCCAAAGCCA	ATCACTGGGG	AGACCGAGCA 4240
4241	CAGGGAGGAC	CAAGGGGAAG	GGGAGAGAAA	GGAAATAAAG	AACAACGTTA	TTTCTTAAAC	GACTTTCTAT	AGGAGTTGTA 4320
4321	AGAAGGTGCA	CATATTTTTT	TAAATCTCAC	TGGCAATATT	CAAAGTTTTT	ATTGTGTCTT	AACAAAGGTG	TGGTAGACAC 4400
4401	TCTTGAGCTG	GACTTAGATT	TTATTCTTCC	TTGCAGAGTA	GTGTTAGAAT	AGATGGCCTA	CAGAAAAAAA	AGGTTCTGGG 4480
4481	ATCTACATGG	CAGGGAGGGC	TGCACTGACA	TTGATGCCTG	GGGGACCTTT	TGCCTCGACT	CGTGCCGGAA	ATCTGATCGT 4560
4561	AATCAGGGTA	CAGAACTTAC	TAGTTTTGTC	TAGGAGTATG	TTGTATGACT	AGGATTTGTG	CTATTATCTC	ATTCAACAAC 4640
4641	ATAGAGCAAG	AATAGTGAGC	TAACTGAGCT	AGACACTCAA	TTAATCCGCT	ACTGGCTTCA	AGTCAGAACT	TTGTCAATTA 4720
4721	TCATCGACTC	CGGGACGGTC	ATATATGTAT	TACATTTCTA	CATTTTTAAT	ACTCACATGG	GCTTATGCAT	TAAGTTTAAAT 4800
4801	TGTGATAAAT	TTGTGCTGGT	CCAGTATATG	CAATACACTT	TAATGGTTTA	TTCTTGTCTAT	AAAAATGTGC	AATATGGAGA 4880
4881	TGTATACAAG	TCTTTACT						4898

FIG. 106

	10	20	30	40	50	60	70	80
1	MEGHVMI AFL	PTILNQLFRV	LTRATQEEVA	VNVTRV I I HV	VAQCHEEGLE	SHLRSYVKYA	YKAEPYVASE	YKTVHEELTK 80
81	SMTTILKPSA	DFLT SNKLLR	YSWFFFDVLI	KSMAQH LIEN	SKVKLLRNQR	FPASYHHAAE	TVVNMLMPHI	TQKFGDNPEA 160
161	SKNANHSLAV	FIKRCFTFMD	RGFVFKQINN	YISCFAPGDP	KTLFEYKFEF	LRVVCNHEHY	IPLNLPMPFG	KGRIQRYQDL 240
241	QLDYSLTDEF	CRNHFLVGLL	LREVG TALQE	FREVR LIAIS	VLKNLLIKHS	FDDRYASRSH	QARIATLYLP	LPGLLIENVQ 320
321	RINVRDVSPF	PVNAGMTVKD	ESLALPAVNP	LVTPOKGSTL	DNSLHKDLIG	AISGIASPYT	TSTPNINSVR	NADSRGSLIS 400
401	TDSGNSLPER	NSEKSNSLDK	HQQSSTLGNS	VVRCDKLDQS	EIKSLLMCFL	YILKSMSDDA	LFTYWNKAST	SELMOFFTIS 480
481	EVCLHQFOYM	GKRYIARNQE	GLGPIVHDRK	SQTLPVSRNR	TGMMHARLQQ	LGSLDNSLTF	NHSYGHSDAD	VLHQSLLEAN 560
561	IATEVCLTAL	DTLSLFTLAF	KNQLLADHGH	NPLMKKVFDV	YLCFLQKHQS	ETALKNVFTA	LRSLIYKFPS	TFYEGRADMC 640
641	AALCYEILKC	CNSKLSSIRT	EASQLLYFLM	RNNFDYTGKK	SFVRTHLQVI	ISVSQLIADV	VGIGETRFOQ	SLSIINN CAN 720
721	SDRLIKHTSF	SSDVKD LTKR	IRTVLMATAQ	MKEHENDPEM	LVDLQYSLAK	SYASTPELRK	TWLD SMARIH	VKN GDLSEAA 800
801	MCYVHV TALV	AEYLTRKGVF	RQGCTAFRVI	TPNIDEEASM	MEDVGMQDVH	FNEDVLMELL	EQCADGLWKA	ERYELIADIY 880
881	KLIIP IYEKR	RDFFEDEDGK	EYIYKEPKLT	PLSEI SORLL	KLYSDKFGSE	NVKMIQDSGK	VNPKDLDSKY	AYIQVTHVIP 960
961	FFDEKELQER	KTEFERSHNI	RRFMFEMPFT	QTGKRQGGVE	EQCKRR TILT	AIHCFFPYVKK	RIPVMYQHHT	DLNPIEVAID 1040
1041	EMSKKVAELR	QLCSSAEVDM	IKLQ LKQGS	VSVQVNAGPL	AYARAFLDDT	NTKRYPDNKV	KLLKEVFRQF	VEACQALAV 1120
1121	NERLIKEDQL	EYQEEMKANY	REMAKELSEI	MHEQICPLEE	KTSVLPNSLH	IFNAISGTPT	STMVHGMTSS	SSVV 1194

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Fig. 106 (cont.)



	10	20	30	40	50	60	70	80
1	AATTGTAATA	CGACTCACTA	TAGGGCGAAT	TGGGTACCGG	GCCCCCCTC	GAGGTCGACG	GTATCGATAA	GCTTGATATC 80
81	GAATTCGGCA	CGAGTTTAC	ACCATCACCA	AAACCCAGAA	TTTTATGATG	AGATTAAAT	AGAGTTGCC	ACTCAGCTGC 160
161	ATGAAAAGCA	CCACCTGTG	CTCACATTCT	TCCATGTCAG	CTGTGACAAC	TCAAGTAAAG	GAAGCACGAA	GAAGAGGGAT 240
241	GTCGTTGAAA	CCCAAGTTGG	CTACTCCTGG	CTTCCCCTCC	TGAAAGACGG	AAGGGTGGTG	ACAAGCGAGC	AGCACATCCC 320
321	GGTCTCGGCG	AACCTTCCTT	CGGGCTATCT	TGGCTACCAA	GAGCTTGGGA	TGGGCAGGCA	TTATGGTCCG	GAAATTAAAT 400
401	GGGTAGATGG	AGGCAAGCCA	CTGCTGAAAA	TTTCCACTCA	TCTGGTTTCT	ACAGGGATAC	TCAGGATCAG	CATTTACATA 480
481	ATTTTTTCCA	GTACTGTCAG	AAAACCGAAT	CTGGAGCCCA	AGCCTTAGGA	AACGAACCTG	TAAAGTACCT	TAAGAGTCTG 560
561	CATGCGATGG	AAGGCCACGT	GATGATCGCC	TTCTTGCCCA	CTATCCTAAA	CCAGCTGTTC	CGAGTCCTCA	CCAGAGCCAC 640
641	ACAGGAAGAA	GTCGCGGTTA	ACGTGACTCG	GGTCATTATT	CATGTGGTTG	CCCAGTGCCA	TGAGGAAGGA	TTGGAGAGCC 720
721	ACTTGAGGTC	ATATGTTAAG	TACGCGTATA	AGGCTGAGCC	ATATGTTGCC	TCTGAATACA	AGACAGTGCA	TGAAGAACTG 800
801	ACCAAATCCA	TGACCACGAT	TCTCAAGCCT	TCTGCCGATT	TCCTCACCAG	CAACAACTA	CTGAGGTAAT	CATGGTTTTT 880
881	CTTTGATGTA	CTGATCAAAT	CTATGGCTCA	GCATTTGATA	GAGAACTCCA	AAGTTAAGTT	GCTGCGAAAC	CAGAGATTTC 960
961	CTGCATCCTA	TCATCATGCA	GCGGAAACCG	TTGTAAATAT	GCTGATGCCA	CACATCACTC	AGAAGTTTGG	AGATAATCCA 1040
1041	GAGGCATCTA	AGAACGCGAA	TCATAGCCTT	GCTGTCTTCA	TCAAGAGATG	TTTCACTTTC	ATGGACAGGG	GCTTTGTCTT 1120
1121	CAAGCAGATC	AACAACCTAC	TTAGCTGTTT	TGCTCCTGGA	GACCCAAAGA	CCCTCTTTGA	ATACAAGTTT	GAATTTCTCC 1200
1201	GTGTAGTGTG	CAACCATGAA	CATTATATTC	CGTTGAACTT	ACCAATGCCA	TTTGGAAAAG	GCAGGATTCA	AAGATACCAA 1280
1281	GACCTCCAGC	TTGACTACTC	ATTAACAGAT	GAGTCTGCA	GAAACCACTT	CTTGGTGGGA	CTGTTACTGA	GGGAGGTGGG 1360
1361	GACAGCCCTC	CAGGAGTTCC	GGGAGGTCCG	TCTGATCGCC	ATCAGTGTGC	TCAAGAACCT	GCTGATAAAG	CATTCTTTTG 1440
1441	ATGACAGATA	TGCTTCAAGG	AGCCATCAGG	CAAGGATAGC	CACCTCTAC	CTGCCCTGTG	TTGGTCTGCT	GATTGAAAAC 1520
1521	GTCCAGCGGA	TCAATGTGAG	GGATGTGTCA	CCCTTCCCTG	TGAACGCGGG	CATGACCGTG	AAGGATGAAT	CCCTGGCTCT 1600
1601	ACCAGCTGTG	AATCCGCTGG	TGACGCCGCA	GAAGGGAAGC	ACCCTGGACA	ACAGCCTGCA	CAAGGACCTG	CTGGGCGCCA 1680
1681	TCTCCGGCAT	TGCTTCTCCA	TATACAACCT	CAACTCCAAA	CATCAACAGT	GTGAGAAATG	CTGATTGAG	AGGATCTCTC 1760
1761	ATAAGCACAG	ATTCCGGTAA	CAGCCTTCCA	GAAAGGAATA	GTGAGAAGAG	CAATTCCCTG	GATAAGCACC	AACAAAGTAG 1840
1841	CACATTGGGA	AATTCCGTGG	TTGCTGTGTA	TAAACTTGAC	CAGTCTGAGA	TTAAGAGCCT	ACTGATGTGT	TTCTCTTACA 1920
1921	TCTTAAAGAG	CATGTCTGAT	GATGCTTTGT	TTACATATTG	GAACAAGGCT	TCAACATCTG	AACTTATGGA	TTTTTTTACA 2000
2001	ATATCTGAAG	TCTGCCGTGA	CCAGTCCAG	TACATGGGA	AGCGATACAT	AGCCAGGAAC	CAGGAGGGGT	TGGGACCCAT 2080
2081	AGTTCATGAT	CGAAAGTCTC	AGACATTGCC	TGTTTCCCGT	AACAGAACAG	GAATGATGCA	TGCCAGATTG	CAGCAGCTGG 2160
2161	GCAGCCTGGA	TAACTCTCTC	ACTTTTAACC	ACAGCTATGG	CCACTCGGAC	GCAGATGTTT	TGCACCAGTC	ATTACTTGAA 2240
2241	GCCAACATTG	CTACTGAGGT	TGCGCTGACA	GCTCTGGACA	CGCTTTCTCT	ATTTACATTG	GCGTTTAAGA	ACCAGCTCCT 2320
2321	GGCCGACCAT	GGACATAATC	CTCTCATGAA	AAAAGTTTTT	GATGTCTACC	TGTGTTTTCT	TCAAAAACAT	CAGTCTGAAA 2400
2401	CGGCTTTTAA	AAATGTCTTC	ACTGCCTTAA	GGTCTTAAAT	TTATAAGTTT	CCCTCAACAT	TCTATGAAGG	GAGAGCGGAC 2480
2481	ATGTGTGCGG	CTCTGTGTTA	CGAGATTCTC	AAGTGTCTGA	ACTCCAAGCT	GAGCTCCATC	AGGACGGAGG	CCTCCCAGCT 2560
2561	GCTCTACTTC	CTGATGAGGA	ACAACCTTGA	TTACACTGGA	AAGAAGTCCT	TTGTCCGGAC	ACATTTGCAA	GTCATCATAT 2640
2641	CTGTCAGCCA	GCTGATAGCA	GACGTTGTTG	GCATTGGGGA	AACCAGATTG	CAGCAGTCCC	TGTCCATCAT	CAACAACGTG 2720
2721	GCCAACAGTG	ACCGGCTTAT	TAAGCACACC	AGCTTCTCCT	CTGATGTGAA	GGACTTAACC	AAAAGGATAC	GCACGGTGCT 2800
2801	AATGGCCACC	GCCAGATGTA	AGGAGCATGA	GAACGACCCA	GAGATGCTGG	TGGACCTCCA	GTACAGCCTG	GCCAAATCCT 2880
2881	ATGCCAGCAC	GCCCGAGCTC	AGGAAGACGT	GGCTCGACAG	CATGGCCAGG	ATCCATGTCA	AAAATGGCGA	TCTCTCAGAG 2960
2961	GCAGCAATGT	GCTATGTCCA	CGTAACAGCC	CTAGTGGCAG	AATATCTCAC	ACGGAAGGCG	GTGTTTAGAC	AAGGATGCAC 3040
3041	CGCCTTCAGG	GTCATTACCC	CAAACATCGA	CGAGGAGGCC	TCCATGATGG	AAGACGTGGG	GATGCAGGAT	GTCCATTTC 3120
3121	ACGAGGATGT	GCTGATGGAG	CTCCTTGAGC	AGTGCGCAGA	TGGACTCTGG	AAAGCCGAGC	GCTACGAGCT	CATCGCCGAC 3200
3201	ATCTACAAAC	TTATCATCCC	CATTTATGAG	AAGCGGAGGG	ATTTCTTTGA	AGATGAAGAT	GGAAAGGAGT	ATATTTACAA 3280
3281	GGAACCCAAA	CTCACACCGC	TGTCGGAAAT	TTCTCAGAGA	CTCCTTAAAC	TGTAATCGGA	TAAATTTGGT	TCTGAAAATG 3360
3361	TCAAAATGAT	ACAGGATTCT	GGCAAGGTCA	ACCCTAAGGA	TCTGGATTCT	AAGTATGCAT	ACATCCAGGT	GACTCACGTC 3440
3441	ATCCCTTTCT	TTGACGAAAA	AGAGTTGCAA	GAAAGGAAAA	CAGAGTTTGA	GAGATCCAC	AACATCCGCC	GCTTCATGTT 3520
3521	TGAGATGCCA	TTTACGCGA	CCGGGAAGAG	GCAGGGCGGG	GTGGAAGAGC	AGTGCAACCG	GCGCACCATC	CTGACAGCCA 3600
3601	TACACTGCTT	CCCTTATGTG	AAGAAGCGCA	TCCCTGTCAT	GTACCAGCAC	CACACTGACC	TGAACCCCAT	CGAGGTGGCC 3680
3681	ATTGACGAGA	TGAGTAAGAA	GGTGGCGGAG	CTCCGGCAGC	TGTGCTCCTC	GGCCGAGGTG	GACATGATCA	AACTGCAGCT 3760
3761	CAAACCTCCAG	GGCAGCGTGA	GTGTTCAAGT	CAATGCTGGC	CCACTAGCAT	ATGCGCGAGC	TTTCTTAGAT	GATACAAACA 3840
3841	CAAAGCGATA	TCCTGACAAT	AAAGTGAAGC	TGCTTAAGGA	AGTTTTTCAGG	CAATTTGTGG	AAGCTTGCGG	TCAAGCCTTA 3920
3921	GCGGTAAACG	AACGTCTGAT	TAAAGAAGAC	CAGCTCGAGT	ATCAGGAAGA	AATGAAAGCC	AACTACAGGG	AAATGGCGAA 4000
4001	GGAGCTTTCT	GAAATCATGC	ATGAGCAGAT	CTGCCCCCTG	GAGGAGAAGA	CGAGCGTCTT	ACCGAATTCC	CTTCACATCT 4080
4081	TCAACGCCAT	CAGTGGGACT	CCAACAAGCA	CAATGGTTCA	CGGGATGACC	AGCTCGTCTT	CGGTGCTGTG	ATTACATCTC 4160
4161	ATGGCCCGTG	TGTGGGGACT	TGCTTTGTCA	TTTGCAAAC	CAGGATGCTT	TCCAAAGCCA	ATCACTGGGG	AGACCGAGCA 4240
4241	CAGGGAGGAC	CAAGGGGAAG	GGGAGAGAAA	GGAAATAAAG	AACAACGTTA	TTTCTTAAAC	GACTTTCTAT	AGGAGTTGTA 4320
4321	AGAAGGTGCA	CATATTTTCT	TAAATCTCAC	TGGCAATATT	CAAAGTTTTC	ATTGTGTCTT	AACAAAGGTG	TGGTAGACAC 4400
4401	TCTTGAGCTG	GACTTAGATT	TTATTCTTCC	TTGCAGAGTA	GTGTTAGAAT	AGATGGCCTA	CAGAAAAAAA	AGGTTCTGGG 4480
4481	ATCTACATGG	CAGGGAGGGC	TGCACTGACA	TTGATGCCTG	GGGGACCTTT	TGCCTCGACT	CGTGCCGGAA	ATCTGATCGT 4560
4561	AATCAGGGTA	CAGAACCTAC	TAGTTTTGTC	TAGGAGTATG	TTGTATGACT	AGGATTTGTG	CTATTATCTC	ATTCAACAAC 4640
4641	ATAGAGCAAG	AATAGTGAGC	TAACTGAGCT	AGACACTCAA	TTAATCCGCT	ACTGGCTTCA	AGTCAGAACT	TTGTCATTAA 4720
4721	TCATCGACTC	CGGGACGGTC	ATATATGTAT	TACATTTCTA	CATTTTAAAT	ACTCACATGG	GCTTATGCAT	TAAGTTTAAAT 4800
4801	TGTGATAAAT	TTGTGCTGGT	CCAGTATATG	CAATACACTT	TAATGGTTTA	TTCTTGTGCT	AAAAATGTGC	AATATGGAGA 4880
4881	TGTATACAAG	TCTTTACT						4898

FIG. 10H



Exon 1A (-182 to -102)

GCAGGGGAAAAACCTGGCCCCATGATTCACTTACTTCCCACCGGATCTCTCCCATGACACGTGAGGATTA  
TTACAATTTAA -102

Exon 1B (-219 to -102)

TTATCCCTTTACTACTTGCGAAGTGAGTTCGGTAGATGGGAGTGGAGAAGAGAACCTTAGAATCATTGTTTAGTCTTCAT  
CTTTCACAGCTCAGGCTGAAGGCCTTTCCTTGCTGAGA -102

Exon 1C (-143 to -102)

GCGGCAGAGCGTGTCTGAGGTGGTGCGCGGCTCCGTGCTCCT -102

Exon2 and the rest of human CLASP2 cDNA

-101 -79  
GGCAAAGCCAAAGCTAATTGAGC

-78 -1  
AAGCTAATTGAGCCACTCGACTATGAAAATGTCATCGTCCAGAAGAAGACTCAGATCCTGAACGACTGTTTACGGGAG

1/1 31/11  
ATG CTG CTC TTC CCT TAC GAT GAC TTT CAG ACG GCC ATC CTG AGA CGA CAG GGT CGA TAC  
Met leu leu phe pro tyr asp asp phe gln thr ala ile leu arg arg gln gly arg tyr

61/21 91/31  
ATA TGC TCA ACA GTG CCT GCG AAG GCG GAA GAG GAA GCA CAG AGC TTG TTT GTT ACA GAG  
ile cys ser thr val pro ala lys ala glu glu glu ala gln ser leu phe val thr glu

121/41 151/51  
TGC ATC AAA ACC TAT AAC TCT GAC TGG CAT CTT GTG AAC TAT AAA TAT GAA GAT TAC TCA  
cys ile lys thr tyr asn ser asp trp his leu val asn tyr lys tyr glu asp tyr ser

181/61 211/71  
GGA GAG TTT CGA CAG CTT CCG AAC AAA GTG GTC AAG TTG GAT AAA CTT CCA GTT CAT GTC  
gly glu phe arg gln leu pro asn lys val val lys leu asp lys leu pro val his val

241/81 271/91  
TAT GAA GTT GAC GAG GAG GTC GAC AAA GAT GAG GAT GCT GCC TCC CTT GGT TCC CAG AAG  
tyr glu val asp glu glu val asp lys asp glu asp ala ala ser leu gly ser gln lys

301/101 331/111  
GGT GGG ATC ACC AAG CAT GGC TGG CTG TAC AAA GGC AAC ATG AAC AGT GCC ATC AGC GTG  
gly gly ile thr lys his gly trp leu tyr lys gly asn met asn ser ala ile ser val

361/121 391/131  
ACC ATG AGG TCA TTT AAG AGA CGA TTT TTC CAC CTG ATT CAA CTT GGC GAT GGA TCC TAT  
thr met arg ser phe lys arg arg phe phe his leu ile gln leu gly asp gly ser tyr

421/141 451/151  
AAT TTG AAT TTT TAT AAA GAT GAA AAG ATC TCC AAA GAA CCA AAA GGA TCA ATA TTT CTG  
asn leu asn phe tyr lys asp glu lys ile ser lys glu pro lys gly ser ile phe leu

481/161 511/171  
GAT TCC TGT ATG GGT GTC GTT CAG AAC AAC AAA GTC AGG CGT TTT GCT TTT GAG CTC AAG  
asp ser cys met gly val val gln asn asn lys val arg arg phe ala phe glu leu lys

541/181	571/191
ATG CAG GAC AAA AGT AGT TAT CTC TTG GCA GCA GAC AGT GAA GTG GAA ATG GAA GAA TGG	
met gln asp lys ser ser tyr leu leu ala ala asp ser glu val glu met glu glu trp	
601/201	631/211
ATC ACA ATT CTA AAT AAG ATC CTC CAG CTC AAC TTT GAA GCT GCA ATG CAA GAA AAG CGA	
ile thr ile leu asn lys ile leu gln leu asn phe glu ala ala met gln glu lys arg	
661/221	691/231
AAT GGC GAC TCT CAC GAA GAT GAT GAA CAA AGC AAA TTG GAA GGT TCT GGT TCC GGT TTA	
asn gly asp ser his glu asp asp glu gln ser lys leu glu gly ser gly ser gly leu	
721/241	751/251
GAT AGC TAC CTG CCG GAA CTT GCC AAG AGT GCA AGA GAA GCA GAA ATC AAA CTA AAA AGT	
asp ser tyr leu pro glu leu ala lys ser ala arg glu ala glu ile lys leu lys ser	
781/261	811/271
GAA AGC AGA GTC AAA CTT TTT TAT TTG GAC CCA GAT GCC CAG AAG CTT GAC TTC TCA TCA	
glu ser arg val lys leu phe tyr leu asp pro asp ala gln lys leu asp phe ser ser	
841/281	871/291
GCT GAG CCA GAA GTG AAG TCA TTT GAA GAG AAG TTT GGA AAA AGG ATC CTT GTC AAG TGC	
ala glu pro glu val lys ser phe glu glu lys phe gly lys arg ile leu val lys cys	
901/301	931/311
AAT GAT TTA TCT TTC AAT TTG CAA TGC TGT GTT GCC GAA AAT GAA GAA GGA CCC ACT ACA	
asn asp leu ser phe asn leu gln cys cys val ala glu asn glu glu gly pro thr thr	
961/321	991/331
AAT GTT GAA CCT TTC TTT GTT ACT CTA TCC CTG TTT GAC ATA AAA TAC AAC CGG AAG ATT	
asn val glu pro phe phe val thr leu ser leu phe asp ile lys tyr asn arg lys ile	
1021/341	1051/351
TCT GCC GAT TTC CAC GTA GAC CTG AAC CAT TTC TCA GTG AGG CAA ATG CTC GCC ACC ACG	
ser ala asp phe his val asp leu asn his phe ser val arg gln met leu ala thr thr	
1081/361	1111/371
TCC CCG GCG CTG ATG AAT GGC AGT GGG CAG AGC CCA TCT GTC CTC AAG GGC ATC CTT CAT	
ser pro ala leu met asn gly ser gly gln ser pro ser val leu lys gly ile leu his	
1141/381	1171/391
GAA GCC GCC ATG CAG TAT CCG AAG CAG GGA ATA TTT TCA GTC ACT TGT CCT CAT CCA GAT	
glu ala ala met gln tyr pro lys gln gly ile phe ser val thr cys pro his pro asp	
1201/401	1231/411
ATA TTT CTT GTG GCC AGA ATT GAA AAA GTC CTT CAG GGG AGC ATC ACA CAT TGC GCT GAG	
ile phe leu val ala arg ile glu lys val leu gln gly ser ile thr his cys ala glu	
1261/421	1291/431
CCA TAT ATG AAA AGT TCA GAC TCT TCT AAG GTG GCC CAG AAG GTG CTG AAG AAT GCC AAG	
pro tyr met lys ser ser asp ser ser lys val ala gln lys val leu lys asn ala lys	
1321/441	1351/451
CAG GCA TGC CAA AGA CTA GGA CAG TAT AGA ATG CCA TTT GCT TGG GCA GCA AGG ACA TTG	
gln ala cys gln arg leu gly gln tyr arg met pro phe ala trp ala ala arg thr leu	

1381/461	1411/471
TTT AAG GAT GCA TCT GGA AAT CTT GAC AAA AAT GCC AGA TTT TCT GCC ATC TAC AGG CAA	
phe lys asp ala ser gly asn leu asp lys asn ala arg phe ser ala ile tyr arg gln	
1441/481	1471/491
GAC AGC AAT AAG CTA TCC AAT GAT GAC ATG CTC AAG TTA CTT GCA GAC TTT CGG AAA CCT	
asp ser asn lys leu ser asn asp asp met leu lys leu leu ala asp phe arg lys pro	
1501/501	1531/511
GAG AAG ATG GCT AAG CTC CCA GTG ATT TTA GGC AAT CTA GAC ATT ACA ATT GAT AAT GTT	
glu lys met ala lys leu pro val ile leu gly asn leu asp ile thr ile asp asn val	
1561/521	1591/531
TCC TCA GAC TTC CCT AAT TAT GTT AAT TCA TCA TAC ATT CCC ACA AAA CAA TTT GAA ACC	
ser ser asp phe pro asn tyr val asn ser ser tyr ile pro thr lys gln phe glu thr	
1621/541	1651/551
TGC AGT AAA ACT CCC ATC ACG TTT GAA GTG GAG GAA TTT GTG CCC TGC ATA CCA AAA CAC	
cys ser lys thr pro ile thr phe glu val glu glu phe val pro cys ile pro lys his	
1681/561	1711/571
ACT CAG CCT TAC ACC ATC TAC ACC AAT CAC CTT TAC GTT TAT CCT AAG TAC TTG AAA TAC	
thr gln pro tyr thr ile tyr thr asn his leu tyr val tyr pro lys tyr leu lys tyr	
1741/581	1771/591
GAC AGT CAG AAG TCT TTT GCC AAG GCT AGA AAT ATT GCG ATT TGC ATT GAA TTC AAA GAT	
asp ser gln lys ser phe ala lys ala arg asn ile ala ile cys ile glu phe lys asp	
1801/601	1831/611
TCA GAT GAG GAA GAC TCT CAG CCC CTT AAG TGC ATT TAT GGC AGA CCT GGT GGG CCA GTT	
ser asp glu glu asp ser gln pro leu lys cys ile tyr gly arg pro gly gly pro val	
1861/621	1891/631
TTC ACA AGA AGC GCC TTT GCT GCA GTT TTA CAC CAT CAC CAA AAC CCA GAA TTT TAT GAT	
phe thr arg ser ala phe ala ala val leu his his his gln asn pro glu phe tyr asp	
1921/641	1951/651
GAG ATT AAA ATA GAG TTG CCC ACT CAG CTG CAT GAA AAG CAC CAC CTG TTG CTC ACA TTC	
glu ile lys ile glu leu pro thr gln leu his glu lys his his leu leu leu thr phe	
1981/661	2011/671
TTC CAT GTC AGC TGT GAC AAC TCA AGT AAA GGA AGC ACG AAG AAG AGG GAT GTC GTT GAA	
phe his val ser cys asp asn ser ser lys gly ser thr lys lys arg asp val val glu	
2041/681	2071/691
ACC CAA GTT GGC TAC TCC TGG CTT CCC CTC CTG AAA GAC GGA AGG GTG GTG ACA AGC GAG	
thr gln val gly tyr ser trp leu pro leu leu lys asp gly arg val val thr ser glu	
2101/701	2131/711
CAG CAC ATC CCG GTC TCG GCG TAC CTT CCT TCG GGC CAT CTT GGC TAC CAA GAG CTT GGG	
gln his ile pro val ser ala tyr leu pro ser gly his leu gly tyr gln glu leu gly	
2161/721	2191/731
ATG GGC AGG CAT TAT GGT CCG GAA ATT AAA TGG GTA GAT GGA GGC AAG CCA CTG CTG AAA	
met gly arg his tyr gly pro glu ile lys trp val asp gly gly lys pro leu leu lys	



2221/741	2251/751
ATT TCC ACT CAT CTG GTT TCT ACA GTG TAT	ACT CAG GAT CAG CAT TTA CAT AAT TTT TTC
ile ser thr his leu val ser thr val tyr	thr gln asp gln his leu his asn phe phe
2281/761	2311/771
CAG TAC TGT CAG AAA ACC GAA TCT GGA GCC	CAA GCC TTA GGA AAC GAA CTT GTA AAG TAC
gln tyr cys gln lys thr glu ser gly ala	gln ala leu gly asn glu leu val lys tyr
2341/781	2371/791
CTT AAG AGT CTG CAT GCG ATG GAA GGC CAC	GTG ATG ATC GCC TTC TTG CCC ACT ATC CTA
leu lys ser leu his ala met glu gly his	val met ile ala phe leu pro thr ile leu
2401/801	2431/811
AAC CAG CTG TTC CGA GTC CTC ACC AGA GCC	ACA CAG GAA GAA GTC GCG GTT AAC GTG ACT
asn gln leu phe arg val leu thr arg ala	thr gln glu glu val ala val asn val thr
2461/821	2491/831
CGG GTC ATT ATT CAT GTG GTT GCC CAG TGC	CAT GAG GAA GGA TTG GAG AGC CAC TTG AGG
arg val ile ile his val val ala gln cys	his glu glu gly leu glu ser his leu arg
2521/841	2551/851
TCA TAT GTT AAG TAC GCG TAT AAG GCT GAG	CCA TAT GTT GCC TCT GAA TAC AAG ACA GTG
ser tyr val lys tyr ala tyr lys ala glu	pro tyr val ala ser glu tyr lys thr val
2581/861	2611/871
CAT GAA GAA CTG ACC AAA TCC ATG ACC ACG	ATT CTC AAG CCT TCT GCC GAT TTC CTC ACC
his glu glu leu thr lys ser met thr thr	ile leu lys pro ser ala asp phe leu thr
2641/881	2671/891
AGC AAC AAA CTA CTG AGG TAC TCA TGG TTT	TTC TTT GAT GTA CTG ATC AAA TCT ATG GCT
ser asn lys leu leu arg tyr ser trp phe	phe phe asp val leu ile lys ser met ala
2701/901	2731/911
CAG CAT TTG ATA GAG AAC TCC AAA GTT AAG	TTG CTG CGA AAC CAG AGA TTT CCT GCA TCC
gln his leu ile glu asn ser lys val lys	leu leu arg asn gln arg phe pro ala ser
2761/921	2791/931
TAT CAT CAT GCA GCG GAA ACC GTT GTA AAT	ATG CTG ATG CCA CAC ATC ACT CAG AAG TTT
tyr his his ala ala glu thr val val asn	met leu met pro his ile thr gln lys phe
2821/941	2851/951
GGA GAT AAT CCA GAG GCA TCT AAG AAC GCG	AAT CAT AGC CTT GCT GTC TTC ATC AAG AGA
gly asp asn pro glu ala ser lys asn ala	asn his ser leu ala val phe ile lys arg
2881/961	2911/971
TGT TTC ACC TTC ATG GAC AGG GGC TTT GTC	TTC AAG CAG ATC AAC AAC TAC ATT AGC TGT
cys phe thr phe met asp arg gly phe val	phe lys gln ile asn asn tyr ile ser cys
2941/981	2971/991
TTT GCT CCT GGA GAC CCA AAG ACC CTC TTT	GAA TAC AAG TTT GAA TTT CTC CGT GTA GTG
phe ala pro gly asp pro lys thr leu phe	glu tyr lys phe glu phe leu arg val val
3001/1001	3031/1011
TGC AAC CAT GAA CAT TAT ATT CCG TTG AAC	TTA CCA ATG CCA TTT GGA AAA GGC AGG ATT
cys asn his glu his tyr ile pro leu asn	leu pro met pro phe gly lys gly arg ile

3061/1021	3091/1031
CAA AGA TAC CAA GAC CTC CAG CTT GAC TAC TCA TTA ACA GAT GAG TTC TGC AGA AAC CAC	
gln arg tyr gln asp leu gln leu asp tyr ser leu thr asp glu phe cys arg asn his	
3121/1041	3151/1051
TTC TTG GTG GGA CTG TTA CTG AGG GAG GTG GGG ACA GCC CTC CAG GAG TTC CGG GAG GTC	
phe leu val gly leu leu leu arg glu val gly thr ala leu gln glu phe arg glu val	
3181/1061	3211/1071
CGT CTG ATC GCC ATC AGT GTG CTC AAG AAC CTG CTG ATA AAG CAT TCT TTT GAT GAC AGA	
arg leu ile ala ile ser val leu lys asn leu leu ile lys his ser phe asp asp arg	
3241/1081	3271/1091
TAT GCT TCA AGG AGC CAT CAG GCA AGG ATA GCC ACC CTC TAC CTG CCT CTG TTT GGT CTG	
tyr ala ser arg ser his gln ala arg ile ala thr leu tyr leu pro leu phe gly leu	
3301/1101	3331/1111
CTG ATT GAA AAC GTC CAG CGG ATC AAT GTG AGG GAT GTG TCA CCC TTC CCT GTG AAC GCG	
leu ile glu asn val gln arg ile asn val arg asp val ser pro phe pro val asn ala	
3361/1121	3391/1131
GGC ATG ACC GTG AAG GAT GAA TCC CTG GCT CTA CCA GCT GTG AAT CCG CTG GTG ACG CCG	
gly met thr val lys asp glu ser leu ala leu pro ala val asn pro leu val thr pro	
3421/1141	3451/1151
CAG AAG GGA AGC ACC CTG GAC AAC AGC CTG CAC AAG GAC CTG CTG GGC GCC ATC TCC GGC	
gln lys gly ser thr leu asp asn ser leu his lys asp leu leu gly ala ile ser gly	
3481/1161	3511/1171
ATT GCT TCT CCA TAT ACA ACC TCA ACT CCA AAC ATC AAC AGT GTG AGA AAT GCT GAT TCG	
ile ala ser pro tyr thr thr ser thr pro asn ile asn ser val arg asn ala asp ser	
3541/1181	3571/1191
AGA GGA TCT CTC ATA AGC ACA GAT TCG GGT AAC AGC CTT CCA GAA AGG AAT AGT GAG AAG	
arg gly ser leu ile ser thr asp ser gly asn ser leu pro glu arg asn ser glu lys	
3601/1201	3631/1211
AGC AAT TCC CTG GAT AAG CAC CAA CAA AGT AGC ACA TTG GGA AAT TCC GTG GTT CGC TGT	
ser asn ser leu asp lys his gln gln ser ser thr leu gly asn ser val val arg cys	
3661/1221	3691/1231
GAT AAA CTT GAC CAG TCT GAG ATT AAG AGC CTA CTG ATG TGT TTC CTC TAC ATC TTA AAG	
asp lys leu asp gln ser glu ile lys ser leu leu met cys phe leu tyr ile leu lys	
3721/1241	3751/1251
AGC ATG TCT GAT GAT GCT TTG TTT ACA TAT TGG AAC AAG GCT TCA ACA TCT GAA CTT ATG	
ser met ser asp asp ala leu phe thr tyr trp asn lys ala ser thr ser glu leu met	
3781/1261	3811/1271
GAT TTT TTT ACA ATA TCT GAA GTC TGC CTG CAC CAG TTC CAG TAC ATG GGG AAG CGA TAC	
asp phe phe thr ile ser glu val cys leu his gln phe gln tyr met gly lys arg tyr	
3841/1281	3871/1291
ATA GCC AGG AAC CAG GAG GGG TTG GGA CCC ATA GTT CAT GAT CGA AAG TCT CAG ACA TTG	
ile ala arg asn gln glu gly leu gly pro ile val his asp arg lys ser gln thr leu	

3901/1301	3931/1311
CCT GTT TCC CGT AAC AGA ACA GGA ATG ATG CAT GCC AGA TTG CAG CAG CTG GGC AGC CTG	
pro val ser arg asn arg thr gly met met his ala arg leu gln gln leu gly ser leu	
3961/1321	3991/1331
GAT AAC TCT CTC ACT TTT AAC CAC AGC TAT GGC CAC TCG GAC GCA GAT GTT CTG CAC CAG	
asp asn ser leu thr phe asn his ser tyr gly his ser asp ala asp val leu his gln	
4021/1341	4051/1351
TCA TTA CTT GAA GCC AAC ATT GCT ACT GAG GTT TGC CTG ACA GCT CTG GAC ACG CTT TCT	
ser leu leu glu ala asn ile ala thr glu val cys leu thr ala leu asp thr leu ser	
4081/1361	4111/1371
CTA TTT ACA TTG GCG TTT AAG AAC CAG CTC CTG GCC GAC CAT GGA CAT AAT CCT CTC ATG	
leu phe thr leu ala phe lys asn gln leu leu ala asp his gly his asn pro leu met	
4141/1381	4171/1391
AAA AAA GTT TTT GAT GTC TAC CTG TGT TTT CTT CAA AAA CAT CAG TCT GAA ACG GCT TTA	
lys lys val phe asp val tyr leu cys phe leu gln lys his gln ser glu thr ala leu	
4201/1401	4231/1411
AAA AAT GTC TTC ACT GCC TTA AGG TCC TTA ATT TAT AAG TTT CCC TCA ACA TTC TAT GAA	
lys asn val phe thr ala leu arg ser leu ile tyr lys phe pro ser thr phe tyr glu	
4261/1421	4291/1431
GGG AGA GCG GAC ATG TGT GCG GCT CTG TGT TAC GAG ATT CTC AAG TGC TGT AAC TCC AAG	
gly arg ala asp met cys ala ala leu cys tyr glu ile leu lys cys cys asn ser lys	
4321/1441	4351/1451
CTG AGC TCC ATC AGG ACG GAG GCC TCC CAG CTG CTC TAC TTC CTG ATG AGG AAC AAC TTT	
leu ser ser ile arg thr glu ala ser gln leu leu tyr phe leu met arg asn asn phe	
4381/1461	4411/1471
GAT TAC ACT GGA AAG AAG TCC TTT GTC CGG ACA CAT TTG CAA GTC ATC ATA TCT GTC AGC	
asp tyr thr gly lys lys ser phe val arg thr his leu gln val ile ile ser val ser	
4441/1481	4471/1491
CAG CTG ATA GCA GAC GTT GTT GGC ATT GGG GAA ACC AGA TTC CAG CAG TCC CTG TCC ATC	
gln leu ile ala asp val val gly ile gly glu thr arg phe gln gln ser leu ser ile	
4501/1501	4531/1511
ATC AAC AAC TGT GCC AAC AGT GAC CGG CTT ATT AAG CAC ACC AGC TTC TCC TCT GAT GTG	
ile asn asn cys ala asn ser asp arg leu ile lys his thr ser phe ser ser asp val	
4561/1521	4591/1531
AAG GAC TTA ACC AAA AGG ATA CGC ACG GTG CTA ATG GCC ACC GCC CAG ATG AAG GAG CAT	
lys asp leu thr lys arg ile arg thr val leu met ala thr ala gln met lys glu his	
4621/1541	4651/1551
GAG AAC GAC CCA GAG ATG CTG GTG GAC CTC CAG TAC AGC CTG GCC AAA TCC TAT GCC AGC	
glu asn asp pro glu met leu val asp leu gln tyr ser leu ala lys ser tyr ala ser	
4681/1561	4711/1571
ACG CCC GAG CTC AGG AAG ACG TGG CTC GAC AGC ATG GCC AGG ATC CAT GTC AAA AAT GGC	
thr pro glu leu arg lys thr trp leu asp ser met ala arg ile his val lys asn gly	

4741/1581 4771/1591  
GAT CTC TCA GAG GCA GCA ATG TGC TAT GTC CAC GTA ACA GCC CTA GTG GCA GAA TAT CTC  
asp leu ser glu ala ala met cys tyr val his val thr ala leu val ala glu tyr leu

4801/1601 4831/1611  
ACA CGG AAA GGC GTG TTT AGA CAA GGA TGC ACC GCC TTC AGG GTC ATT ACC CCA AAC ATC  
thr arg lys gly val phe arg gln gly cys thr ala phe arg val ile thr pro asn ile

4861/1621 4891/1631  
GAC GAG GAG GCC TCC ATG ATG GAA GAC GTG GGG ATG CAG GAT GTC CAT TTC AAC GAG GAT  
asp glu glu ala ser met met glu asp val gly met gln asp val his phe asn glu asp

4921/1641 4951/1651  
GTG CTG ATG GAG CTC CTT GAG CAG TGC GCA GAT GGA CTC TGG AAA GCC GAG CGC TAC GAG  
val leu met glu leu leu glu gln cys ala asp gly leu trp lys ala glu arg tyr glu

4981/1661 5011/1671  
CTC ATC GCC GAC ATC TAC AAA CTT ATC ATC CCC ATT TAT GAG AAG CGG AGG GAT TTC TTT  
leu ile ala asp ile tyr lys leu ile ile pro ile tyr glu lys arg arg asp phe phe

5041/1681 5071/1691  
GAA GAT GAA GAT GGA AAG GAG TAT ATT TAC AAG GAA CCC AAA CTC ACA CCG CTG TCG GAA  
glu asp glu asp gly lys glu tyr ile tyr lys glu pro lys leu thr pro leu ser glu

5101/1701 5131/1711  
ATT TCT CAG AGA CTC CTT AAA CTG TAC TCG GAT AAA TTT GGT TCT GAA AAT GTC AAA ATG  
ile ser gln arg leu leu lys leu tyr ser asp lys phe gly ser glu asn val lys met

5161/1721 5191/1731  
ATA CAG GAT TCT GGC AAG GTC AAC CCT AAG GAT CTG GAT TCT AAG TAT GCA TAC ATC CAG  
ile gln asp ser gly lys val asn pro lys asp leu asp ser lys tyr ala tyr ile gln

5221/1741 5251/1751  
GTG ACT CAC GTC ATC CCC TTC TTT GAC GAA AAA GAG TTG CAA GAA AGG AAA ACA GAG TTT  
val thr his val ile pro phe phe asp glu lys glu leu gln glu arg lys thr glu phe

5281/1761 5311/1771  
GAG AGA TCC CAC AAC ATC CGC CGC TTC ATG TTT GAG ATG CCA TTT ACG CAG ACC GGG AAG  
glu arg ser his asn ile arg arg phe met phe glu met pro phe thr gln thr gly lys

5341/1781 5371/1791  
AGG CAG GGC GGG GTG GAA GAG CAG TGC AAA CGG CGC ACC ATC CTG ACA GCC ATA CAC TGC  
arg gln gly gly val glu glu gln cys lys arg arg thr ile leu thr ala ile his cys

5401/1801 5431/1811  
TTC CCT TAT GTG AAG AAG CGC ATC CCT GTC ATG TAC CAG CAC CAC ACT GAC CTG AAC CCC  
phe pro tyr val lys lys arg ile pro val met tyr gln his his thr asp leu asn pro

5461/1821 5491/1831  
ATC GAG GTG GCC ATT GAC GAG ATG AGT AAG AAG GTG GCG GAG CTC CGG CAG CTG TGC TCC  
ile glu val ala ile asp glu met ser lys lys val ala glu leu arg gln leu cys ser

5521/1841 5551/1851  
TCG GCC GAG GTG GAC ATG ATC AAA CTG CAG CTC AAA CTC CAG GGC AGC GTG AGT GTT CAG  
ser ala glu val asp met ile lys leu gln leu lys leu gln gly ser val ser val gln

5581/1861	5611/1871
GTC AAT GCT GGC CCA CTA GCA TAT GCG CGA	GCT TTC TTA GAT GAT ACA AAC ACA AAG CGA
val asn ala gly pro leu ala tyr ala arg	ala phe leu asp asp thr asn thr lys arg
5641/1881	5671/1891
TAT CCT GAC AAT AAA GTG AAG CTG CTT AAG	GAA GTT TTC AGG CAA TTT GTG GAA GCT TGC
tyr pro asp asn lys val lys leu leu lys	glu val phe arg gln phe val glu ala cys
5701/1901	5731/1911
GGT CAA GCC TTA GCG GTA AAC GAA CGT CTG	ATT AAA GAA GAC CAG CTC GAG TAT CAG GAA
gly gln ala leu ala val asn glu arg leu	ile lys glu asp gln leu glu tyr gln glu
5761/1921	5791/1931
GAA ATG AAA GCC AAC TAC AGG GAA ATG GCG	AAG GAG CTT TCT GAA ATC ATG CAT GAG CAG
glu met lys ala asn tyr arg glu met ala	lys glu leu ser glu ile met his glu gln
5821/1941	5851/1951
ATC TGC CCC CTG GAG GAG AAG ACG AGC GTC	TTA CCG AAT TCC CTT CAC ATC TTC AAC GCC
ile cys pro leu glu glu lys thr ser val	leu pro asn ser leu his ile phe asn ala
5881/1961	5911/1971
ATC AGT GGG ACT CCA ACA AGC ACA ATG GTT	CAC GGG ATG ACC AGC TCG TCT TCG GTC GTG
ile ser gly thr pro thr ser thr met val	his gly met thr ser ser ser ser val val
5941/1981	5971
TGA TTA CAT CTC ATG GCC CGT GTG TGG GGA	CTT GCT TTG TCA TTT GCA AAC TCA GGA TGC
CAA	
6001	6031
TTT CCA AAG CCA ATC ACT GGG GAG ACC GAG	CAC AGG GAG GAC CAA GGG GAA GGG GAG AGA
6061	6091
AAG GAA ATA AAG AAC AAC GTT ATT TCT TAA	CAG ACT TTC TAT AGG AGT TGT AAG AAG GTG
6121	6151
CAC ATA TTT TTT TAA ATC TCA CTG GCA ATA	TTC AAA GTT TTC ATT GTG TCT TAA CAA AGG
6181	6211
TGT GGT AGA CAC TCT TGA GCT GGA CTT AGA	TTT TAT TCT TCC TTG CAG AGT AGT GTT AGA
6241	6271
ATA GAT GGC CTA CAG AAA AAA AAG GTT CTG	GGA TCT ACA TGG CAG GGA GGG CTG CAC TGA
6301	6331
CAT TGA TGC CTG GGG GAC CTT TTG CCT CGA	CTC GTG CCG GAA ATC TGA TCG TAA TCA GGG
6361	6391
TAC AGA ACT TAC TAG TTT TGT CTA GGA GTA	TGT TGT ATG ACT AGG ATT TGT GCT ATT ATC
6421	6451
TCA TTC AAC AAC ATA GAG CAA GAA TAG TGA	GCT AAC TGA GCT AGA CAC TCA ATT AAT CCG
6481	6511
CTA CTG GCT TCA AGT CAG AAC TTT GTC ATT	AAT CAT CGA CTC CGG GAC GGT CAT ATA TGT
6541	6571
ATT ACA TTT CTA CAT TTT TAA TAC TCA CAT	GGG CTT ATG CAT TAA GTT TAA TTG TGA TAA
6601	6631
ATT TGT GCT GGT CCA GTA TAT GCA ATA CAC	TTT AAT GGT TTA TTC TTG TCA TAA AAA TGT
6661	
GCA ATA TGG AGA TGT ATA CAA GTC TTT ACT	



A. Allelic variations: single nucleotide changes (polymorphism) between CLASP-2 cDNA isoforms

Isoform	Difference	Nucleotide(s)	Consequence
1	polymorphism	862	A to G change; mis-sense mutation
2	polymorphism		A to C change; mis-sense mutation changing codon from histidine to proline
3	polymorphism	2210	A to G change; mis-sense mutation changing codon from asparagine to glutamic acid
4	polymorphism	2225	C to T change; mis-sense mutation changing codon from histidine to tyrosine

B. Alternative splices

Isoform	Difference	Nucleotide(s)	Consequence
1	exon deletion	209-291	premature, in-frame stop codon leading to the production of a truncated, most likely soluble protein

These differences may be found separately or together in various combinations in the different human CLASP-2 isoforms

FIG. 11B

human CLASP-2

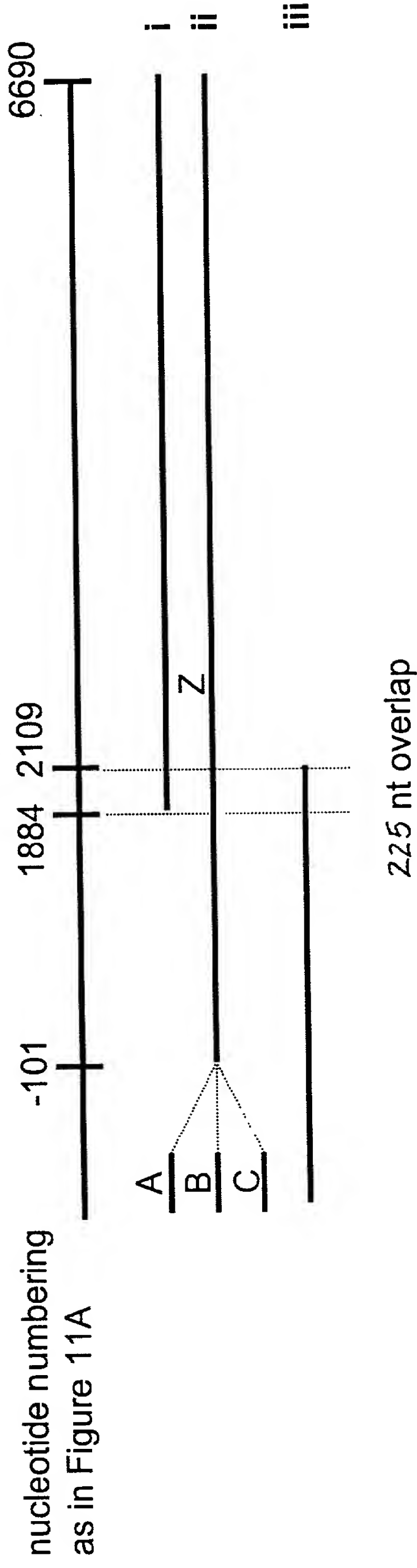


FIG. 11C

1st exon (nucleotides 335 to 445)

TGTCTTGCTTATCTTTTCGCCCTCCAGGCAAAGCCAAAGCTAATTGAGCCACT  
CGACTATGAAAATGTCATCGTCCAGAAGAAGACTCAGATCCTGAACGACTGT  
TTACGGGAGATGCTGCTCTTCCCTTACGATGACTTTCAGGTAAGTAACGTTAT  
GTTTCTATCCGTAGAACCACG

2nd exon (nucleotides 7101-7190)

TTACCCAAGGCTTTTCCTCCTGTTTTTGTTCAGACGGCCATCCTGAGACGA  
CAGGGTCGATACATATGCTCAACAGTGCCTGCGAAGGCGGAAGAGGAAGCA  
CAGAGCTTGTTTGTTACAGAGGTAAGGCTCTTTCCTGCATTAATTTACATTTT  
GAAGTCATTTTCCCCTAACTGCCTCC

3rd exon (nucleotides 11439 to 11521)

TTTTCTATTTTTTAAAATCCCCCTTCAATAGTGCATCAAAACCTATAACTCTGAC  
TGGCATCTTGTGAACATAAATATGAAGATTACTCAGGAGAGTTTCGACAGC  
TTCCGAAGTGAGTAAGCTATATTATACACATAGGGAAAAGTCTTT

4th exon (nucleotides 13987 to 14056)

CTAAAACAAATTTTCTTTGTTGTTTTTATAGCAAAGTGGTCAAGTTGGATAAA  
CTTCCAGTTCATGTCTATGAAGTTGACGAGGAGGTCGACAAAGATGAGGTGG  
GATACCTGCTTGCTGTTGCTTCTCTTTTCACTCTAGATTAA

5th exon (nucleotides 15212 to 15307)

GGAGGTTGACTGCTGGTGTTCCTTCTCTCCTAGGATGCTGCCTCCCTTGGTT  
CCCAGAAGGGTGGGATCACCAAGCATGGCTGGCTGTACAAAGGCAACATGA  
ACAGTGCCATCAGCGTGACCATGAGGGTGAGGACGCACATCACTTTGCCCTC  
CCCTCTCACAAGCCCTTTC

6th exon (nucleotides 16269 to 16404)

TGAAAGAATAGCTGTGTGTATATTTTTCTCTCAGTCATTTAAGAGACGATTTT  
TCCACCTGATTCAACTTGGCGATGGATCCTATAATTTGAATTTTTATAAAGAT  
GAAAAGATCTCCAAAGAACCAAAAGGATCAATATTTCTGGATTCTGTATGG  
GTGTCGTTTCAGGTAAATATGAAAAGAGTTTTACCATTATGTTTTCTTA

7th exon (nucleotides 19459 to 19633)

AAGTATGTCTGTTTATCCTTTTTTTCATTTTCAGAACAAACAAAGTCAGGCGTTTT  
GCTTTTGAGCTCAAGATGCAGGACAAAAGTAGTTATCTCTTGGCAGCAGACA  
GTGAAGTGGAAATGGAAGAATGGATCACAATTCTAAATAAGATCCTCCAGCT  
CAACTTTGAAGCTGCAATGCAAGAAAAGCGAAATGGCGACTCTCACGAAGGT  
AGATAGGCTTGGCTTCCCCCAGGCACATACACACTCT

8th exon (nucleotides 20567 to 20634)

ATTACAAGTGATTCCGATAATCTGTTTTGCCATTTTAGATGATGAACAAAGCA  
AATTGGAAGGTTCTGGTTCCGGTTTAGATAGCTACCTGCCGGAAGTTGCCAAG  
GTAACATCGTCTTATATCTTCTGCTCTTCGTTGAATGC

9th exon (nucleotides 30257 to 30331)

GATTGTGTTAAATGTAATTTTCATGTATCTTGTTATCAGAGTGCAAGAGAAGC  
AGAAATCAAACATAAAAAGTGAAAGCAGAGTCAAACCTTTTTATTGGACCCA  
GATGCCCAGGTAAGAACTATCTAAATGTTTAATATTAAACCAAAT

10th exon (nucleotides 31851 to 31991)

CATAACTTATTTATATGTTTACATTTTCTTTTAAAGAAGCTTGACTTCTCATCA  
GCTGAGCCAGAAAGTGAAGTCATTTGAAGAGAAGTTTGAAAAAGGATCCTTG  
TCAAGTGCAATGATTTATCTTTCAATTTGCAATGCTGTGTTGCCGAAAATGAA  
GAAGGACCCACTACAAATGTAATTTTTCATTTTAAAAATAAACATTAAAAAA  
AAAATAGGCAG

11th exon (nucleotides 32472 to 32675)

CCATGGTGATCATTGGATTGTTTTGTTTTGTTTCAGGTTGAACCTTTCTTTGTTA  
CTCTATCCCTGTTTGACATAAAATACAACCGGAAGATTTCTGCCGATTTCCAC  
GTAGACCTGAACCATTTCTCAGTGAGGCAAATGCTCGCCACCACGTCCCCGG  
CGCTGATGAATGGCAGTGGGCAGAGCCCATCTGTCCTCAAGGGCATCCTTCA  
TGAAGCCGCCATGCAGTATCCGAAGCAGGTGGGGAGTATGAGCCAGCATTC  
CCACTACTCAGACTCACTTTGCATGC

12th exon (nucleotides 33063 to 33185)

GAATTCTGCTTACTGAAGAAAATTGTTTGCCTCCTAGGGAATATTTTCAGTCA  
CTTGTCCTCATCCAGATATATTCTTGTGGCCAGAATTGAAAAAGTCCTTCAG  
GGGAGCATCACACATTGCGCTGAGCCATATATGAAAAGTTCAGACTCTTCTA  
AGGTATGAATGGCTTTTACGCTTTGGGGTGGTAAAAAGCAATCTGAA

13th exon (nucleotides 36702 to 36784)

CAGTATCTCATAGCTTTATTCTCATGTCTTCAAGGTGGCCCAGAAGGTGCTGA  
AGAATGCCAAGCAGGCATGCCAAAGACTAGGACAGTATAGAATGCCATTTGC  
TTGGGCAGCAAGGTAAGGAACACCTTTTATACCTTTTAAATCGATATAGATA  
GGTGCATGG

14th partial exon (nucleotides 37353 to 37475)

GAAACCCAGTTTAGAAATGTTGCTTTGCCATTTTCAGGACATTGTTTAAGGATG  
CATCTGGAAATCTTGACAAAAATGCCAGATTTTCTGCCATCTACAGGCAAGA  
CAGCAATAAGCTATCCAATGATGACATGCTCAAGTTACTTGCAGACTTTCGG  
AA

1 TACCAAGGGCAACTCTGGCACACCCTAAAGTCTGGAAAGGGGACATAGCTAGTCAGGGATGACCCGAGAAATGACTGGAAGCTCCACCAGAA  
93 TGCAGAGCTTCCTTTGTGCTTAAATAACTGAACAAGCATCACTCTGTGTAGCAGGACACCACCCAGCATTTCCTTTGTCCCTTTGGAAACAACT  
185 CTTATTTCTGTTTCTTTGTGATACCAAACTAGCATACTCTAATTGTAGAAAATACAAAACATAGAGTAGAACATACTAAGTTCTTTATCTT  
277 AAGAAATGGCATTGTGTATGAGAATGTCTTGCTTATCTTTTCGCCCTCCAGGCAAAGCCAAAGCTAATTGAGCCACTCGACTATGAAAATG  
369 TCATCGTCCAGAAGAAGACTCAGATCCTGAACGACTGTTTACGGGAGATGCTGCTCTTCCCTTACGATGACTTTCAGGTAAGTAACGTTATG  
461 TTTCTATCCGTAGAACCACGTGTTTGATCTTAACAAGCAGTATTTTCTATGTATTGATTATTGTTTGGTTAGTTAATTATTATTATTATT  
553 ATTTTTTTTGTAGACACAGTCTTGCTCTGTACGCAGGCTTGAGTGCAGTGGTGCCATCTTGGCTCAACGGCAACCTCCGCCTCCTGGGTCA  
645 AGCATTTCTCCTGCCTCAGCCTCTCAAGTAACTGGGATTACAGGCGTGTGCCACCATGCCTGGCTAATTTTTGTCTTTGTATTAGAGACAGG  
737 GTTTTGGCCACGGTGGCCAGGTCGTCCCAAACCTTGGCCTTAAGTGATCTATCTGCCTTGGCCTCTCAAATGTGGGATTATAGGCATGA  
829 GCCACTGTGCCCCGGCCTAATTATGGTTTTTAAAGATGAAAATAAGATGTTATTTAAGAAAGAAAAGTTATTTTATATTCTTCCAAGCATCC  
921 TTCATGAGTTGATAATTTTTAATGGTATTATTTTTGTCATATTAATTATAAGTATGCCAAAATGTGTGTGTGTGTGTGTGTGTGTGTGTGT  
1013 GTGTATCTTGAATAAAAGTGCTATACTCTGTCTGGGCTGATTTAGTGGGCACAAGTGCCCTTCTGTCTTTTGTAGATTGTTTTGATTAGAT  
1105 TTTTTGGCCAAGTACCTTAGAATCTTAATGATGGGCTTGTCTGGGGAGCAGTGGGAGATTTCTGCATGCCTTTTTTTAGGATGGCATTTGGG  
1197 AGCCTGCCTTCAGAGGCCCTGCACCTCTGTGTGGGCTCCAGCAAAGCGTTCAAGGTTAGCCAAGAATGGCCTGAAGTTTACCTCTGTAGTGT  
1289 AATGTGGGTGCTGTTCTTGGAGAAATGTGGAGGACTCAGCACAGCTCCTGCTGTGTGCCGCTCTTTCAGGCTATGGCCTGTGGTTAAGAG  
1381 ACTAACAAGAAGCTGTGAGGCTGTTGAGGAATGAGAATGACATTCTTCTCCAGGAAACCCGGTGGTGTAAATGCCTTGACGCGAGCCACC  
1473 TTGGTCCATTGGAGGTTCTGGTTACTTTCTTGGCTCTCCTGGGACCTGATCCTGGCACTTCTTCTCCTCTCCTTTGATGCTCTTAGTTGG  
1565 ACACCTCTCTACCTGATGCTGTTACCATTAAAGCCCTGTCTTTTGTGAATCGAGCTGCCTTTTTTTTTTAAGCTTCACTGATTCTTTGTGT  
1657 TTGATTCCAAAAGTGTTACATCCATGTACAAAAGATAAATGAGAGGGAAATATTGAAATAAATGACATGAAAAGCCTCCCCACGCCTTCTA  
1749 ATCCCATCCACAGAACATGACTTAACTGTATACAGCTCTGTGCATACTTGTCTTTAGAACTTCCATGTAAATAGAAATTGTTAAATTACG  
1841 ATCCTTGAAGGTTTTTCCACCAAATTTAAGCGACTCCAGCTTACAACAGAGGTGAGAATTTTCACAAATGTTCACTCTTTCTAACTTGT  
1933 AGAGATACCTGGGCCCCAAAATGATTATCTTTAGCTCTGTCTGCATAAAAGGAATGCCATGGGAATGAAATTGACCATTCTGTGGTGT  
2025 TGCTACCAAAGTAACAGGTAAATGGGTTGAGGTATGCCAAACAATACCATGCTTTGCATACTTCATTTTATGACTAACTGCATGGGAACG  
2117 GACTAATAAATGAGAACCTCTGAATGATGCCTTTTGCCTGTGATTGGCAACAATGAAAAGCAAATCAAATGATTATAAATTGTACTGCA  
2209 TGTTGACAAGATTTTCTGTAGTGTGTCTGAGGAAGCTAAAGGTTATCTCAAATTTCTCTCAACATGAAGTATGTGTTCTTCTTGGTATTA  
2301 ATTAAAGTAACAACTTTTTTGAGTTTGCAACCTAGAATGAAAATTCTATTTGTATGACTGAGATAAAATTGCTTAAGAAACAACCAAAGAA  
2393 ACGAGATACAGTTAGTTGAGTGTCTCTTTATCCAGGGAACAGGTATCTGGATGTTTAAAGCAGTTGCAGAATCAGACAGTTTAAACTTTGA  
2485 GAAAACCTCTGTGTCCCCTTTGCTTTTAACTACTCTGGTGATAGCAGGCACAAATATTCTAGGAAAGGCAAAGAACTCACTAGCATTTTGTT  
2577 GGCTAAGGTGATGAGCAAATATTATTTCTGTTTGGGGAGAAGTTTTCTAGAGATTTAGGAGCTTGAATTGGAGCTTTAATCCTCATCACA  
2669 GGAATTGTGATGGGCCCCAGTGAAGTTTGGGTACAATTATTTGTTTTCTTATAGACTCCCCTTTCTTATCAGGTAAAGCCATGTACTCTGT  
2761 GCTTTCTTGTAAATTGTCTCAGTGTATTAACTGTCTAATTAGCTGGATGAGTGAAAGGTCTTAACAGTGCCACAGATTCTTTCTATC  
2853 TGTGTTTTCTTAGGCAGAATAAGAGCAGAATTATTGTATTATTAGAGGCAGAGGGAACAAATTAGATTGGGGAAAGTGTTTTATTTTATATG  
2945 GAAAAGTAATACCAAGTTGGTTAGGAAATGGCAGCAGCAAACGCATGCTGAGGGGTGATTTACTGCCTTAAATAATTTAGCAGTATAAGT  
3037 TAACTATTAAAATAATAGAACTTGGTGTCCATTTCTGCCAAATATATTGAAATGACAATTTACTAAAATATAAGCATGGATAGTGGTGATG  
3129 CTTGTGTACATTTTTCAAGTAGGCACATGTTGATCTTGAGCCTTTACTGGTCAGATCCTAAAGGCATCTACATGTTCTCTAAAATGAGTTG  
3221 TGTCAAGAAAAGATTTGCGGGTTGCATGTAGTTGCCGAGGATGACAGAAGAGTAGTTACTACAACAGCAGCAAAGAAGAGAGACATGAAGT  
3313 AAACGTGGATTTTTTAAAAATCAAAGAATAGGCCAGGCGCACTGGCTCATGCCTGTAATCCCAGCACTTTGGGAGGCCGAGGTGGGCAGATC  
3405 ACAAGGTGAGGAGTTTGAACCTGGGAGGTGGAGTTGCACTGAGCCAAGATCGCGCCATTGCACTCCAGCCTGGGGGACAAGAGGAAAACCTCC  
3497 CCTGTAATCCCAGCTACTCGGGAGGCTGAGGCAGGAGAATCTCTTGAAACCAGGAGACAGAGGTTGCAGTAAGCTGAGATCGTGCCACTGCA  
3589 TTCCAGCCTGGGCGACAGAGTGAGACTCCATCTCAAAACAAACAAAAATCAAATAATAGTTCCCAGCCATCAGGTTATTGATGAAGTAGG  
3681 CTGGGCACGGTGGCTCACACCTGTAATCCCAGCACATTGGGAGTCCGAGGCAGGTGGATCACCTGAGGTGAGGTGTTGAGACCAGCCTGGC  
3773 CAACATGGCAAACCCCGTCTCTACTAAAATAACAAAATTAGCCAGGCATGGTGGTGGGCACCTATAATCCCAGCTACTTTGGGAGGCTGAG  
3865 GCAAGAGAATCGCTTGAACCTGGGAGGTGGAGTTGCACTGAGCCAAGATCGCGCCATTGCACTCCAGCCTGGGGGACAAGAGGAAAACCTCC  
3957 ATCTCAAAAAAAGGGAATATTAAATGAAGTAAAGTACATGTGATCTGCCATGGCCAGGGACAGGAATGCCATGGGGCCTGCAGCCGTCA  
4049 CTAGCTGATGGCCCTTCTTTTTGCAGAATCAGATCCTGTGCTTGGGGATCTCTGCCATCTGTGCTTTGGCTTCATGGTTCTCCTTGCCAGC  
4141 AGCATCTTCTCTTCTAGATCTTTCTACCCTTTAGAGACCCTTGAAATCCCATATTGTCTGAAGCTATTTAAGTCCACAGAACTTTTCCC  
4233 CCCACTGTCTCAATTCTTTTCTACTGCCTGTCTGCACCGTGCACATAAACACTTGAGTATGTGGTCTTGGCTGTTTACGACCTACTTCTTA  
4325 GGCTTCTTGACGCAGGCATCCCGCCCGTGTGTGGTCTGAGAAGGGCTGGCTTTGAGCCTCTGTTCTCCACCCACCTGCCACCTACA  
4417 CATGCACAAAATCCCTTTCTGTAGGTGCTAGGGTTGAATACCCATTGCTTACCTTACTAATAGTAAAATTTTACAAGCATTAGGTTATT  
4509 TTCTTTGATTCATCAAGTAAATATTAATACTGTTTGAACATGTGATAGTCCAGCGACTAGATTTGTAAAATATTTGCAGGATCAATGAT

FIG. 12B  
1 of 10



4601 TTGGTTTGGCAGAAAGTAGGTAATTTCTAAAATTAAAAAATGCAGGTAAACAGGGACTGGAGAGGAGTATTTTTTCCCTAGTGATTAATAAAC  
 4693 CTTTATTTTTCTTATTGTTTTGTTGCTTACCCAGTTTATTTGGCGTAAATCTGAGAACTTACTTTTTCCATGAGCAAAGTTAGAGGTAAAC  
 4785 TTTAACAAGCAGTTAGACAGAGGTAATGACCTTTAGATTAAAAGGTTTTAGGTCAAGCTGTATAAGTTGACTTGTCGCTTAAGACATGATGA  
 4877 GCCTCTGTTTAACTGAAAGTCAAGCCCAGGACGCCTGCCTTTTCCATCAAAGACATGGGATTTGGGTGGCAGCTGACTATTGATTTCCAATG  
 4969 ACGATTCTTCTTCAAGTGGAGGTCTTTTTACCAGATGGTCCTGTTGGTGGGGACATTGTTAACCCCTGCGATTAAACCGACGGCATCTTCATCT  
 5061 GGCTTTTAAGCTCCTTGTATCCTGACTTGTTACACAGCTTACTTATGCTTGTGCGACTATGTAAAGTGACAGTATAATGAGAAAGGTAGTGAG  
 5153 TAGTAAGAATGTTGGGAGACAATTTAAGCTACCATTTCATATTTTCAAAAAATTAGACTTTTGTGTCTGGTGTAAACAAACAGAGGACAGAGC  
 5245 TTGTATGAAAGGATAAAAGAGCGTTAAGGGTTACACGTCCATTAGGATAAAAAAACTAGAATATTTCTTTCTGAAACCTGAAGCCCAGGCCG  
 5337 GGCATGGTGGCTCACGCCTGTAATCTCAGCACTTTGGGAGGTTGAGATGGGAGATTGTTTGAGCCCAGGAGTTTGAGACCAGCCTGGGCAAC  
 5429 ATGGTGAAACCCCATCTCTATTTAAAGAATAAGGCTGGGTGTGGTGGCTCACACCTGTAATCCTAGTGCTTTGGGAGTGTGAGGCAGGTGGA  
 5521 TTGCTTGAGTTCAGGAGTTTGAGACCAGCCTGGGCAACATGGTGAAACCCCATCTGTACTAAAAATACAAAATTAGCCGGGTGTGGTGGCG  
 5613 CCCGCTGTAGTCCAGCTACTCAGGAGGCTGAAGCATGACAATCACTTGAACCTGGGAGGCAGAGGTTGCAGTGAGCCGAGATCATGCCAC  
 5705 TGCATCCAGCCTGGGTGACAGAGAGAGACTCCGTCTCAAAAAAATTAAAAAATTAGGCTGGGCGCAGTGGCTCACGCCTGTAATCCCAGCA  
 5797 CTTTGGGAGGCCGAGGTGGGCAGATCACGAGGTCAGGAGATTGAGACCATCCTGGCTAACACGGTGAAACCCCGTCTCTACTAAAAATACAA  
 5889 AAAATTAGTTGGGCATGGTGGCAGGCGCTGTAGTCCCAGCTGCTCGGGAGGCTGAGGCAGAAGAATGGCGTGAACCCGGGAGGCGGAGCTG  
 5981 GCAGTGAGCTGAGATTGCGCCACTGCACTCCAGACTGGGCGACAGAGCGAGACCTGTCTCAAAAAATAAAATAAAATAAAATAAAATAAAAA  
 6073 ATTAAAAAGAAAAAGAAAAGGAAACCTTAAGCCTAGTTATTGAGGTAGACAGGATGCTACCCCTGCCCTGTCATTTTATTTAAAGAAGCAT  
 6165 TTAAGCCTAATGAACACGAGCAGTTCTAATGTCCGTTGGAGGGGAGGTAGCATTACAGTTCATAGATTCAATTTAGCAATTACTGATTGAGC  
 6257 ATCTTCTGTGTGTCTAGTTATCTATGCTCTTAGGCGCTGGGGATGTGGCAGTGAACAAGAAGAGATGTAAATGACAAGAGATGGATGGTGGT  
 6349 GATGGTTGCACAATTGTGTGAATGTACTTAATGCCACTGAACTGTATACTTAAAAATGTTCAAAATGGCTGGGCATGGTGGCTCACGCCTGT  
 6441 AATCCCAGCACTTTGGGAGGCCGAGGCGGGTGGATCACCTGAGATTAGGAGTTCGAGACCAGCCTGATCAACATGGAGAAACCCCGTCTCTA  
 6533 CTAAAAATACAAAATTAAACGAGCGTGGTGGCGCATGCCTGTAATCCCAGCTACTCGGGAGGCTGAGACAGGAGAATCGCTTGAACCCGGGT  
 6625 GCGCGAGGTTGCAGTGAGCCGAGATCCCGCCATTGCACTCCAGCCTGGGCAACAGAGCAAGACTCCATCTCAAAAAAAAAGTTTCAAATGG  
 6717 TAAATTTATGCATATTTTACCAGAATAAAAAAAGGCAGTTAAGACAAGTAAGATGCTGTGTATGTTGGGGCTAGATCAAGCACTTAGGGGTGGG  
 6809 GTGTTAGGGACTTTGAACGGGGCCTCTACCCCTGCGGGAAGGGCTGAGCTGGAGGGATCTGTGGGCCCCGATCAAGAAAGAAGCAGGAGCTG  
 6901 TAACCCAGCCTGGCTTTGGAACCTGAGGCTGCCAGTGAATCTGTTGTGTGTGACGGAAGGAGGCAGCTGCACTGGATGGGAGAACTGGA  
 6993 GGGACTCTGTGGCTGCCAGGGCCAGCTGCAGGGCACACAGCTGCACCTCTGAGGCTGGCACCTGCCTCCTTCACTTACCCAGGCTTTTCCCTC  
 7085 CTGTTTTTGTTCAGACGGCCATCCTGAGACGACAGGGTCGATACATATGCTCAACAGTGCCTGCGAAGGCGGAAGAGGAAGCACAGAGCT  
 7177 TGTTTGTTACAGAGGTAAGGCTCTTTCTGCAATTAATTTACATTTTGAAGTCATTTTCCCCTAACTGCCTCCTTTTCTTTAAATTTCAAAAT  
 7269 TGTCAAGGAAGTGTCAAAAGGGTAATTGTATTTCTATGATGGAAGTTCAAATAGAATAATGTGAATTTTTCAGACTCTGAACTTGGACAGA  
 7361 AATGTCCACAGGGGCTATTTCTTTTTTACATTTTATTATTTTAAAACCTTTATTTATTTGGAGGGGGCTATATCTGACTACAAAAGTGAA  
 7453 TTCCACAGAATTTATCTCATGGACTTAAAATAAGCAGTAACCTGTAAATGAATTCAGTGGAATCTGTGGGAGGTCTTGTTATTGATACTGT  
 7545 TTTTAAGGGTGACACACACATTTATGTATCATTTATTTTCAATTTATATTTTGAAGTTATTTTGGACTAGTATTCATGAAGTACTA  
 7637 GCTGATAATAAGCAGGGTCTATCGCTAGTCAATATATATTATTATATATATTGATTACTATATATATTCCTAATCAAGATACATTGATTAAT  
 7729 ATTATTTTTGTTTGAAAATGCAATAAAATTATCTTATGGAAGAAAGATAAATTATTTACTTTTTTATTTTTTATTTTTTGGAGACAG  
 7821 AGTCTTGCTCTGTTGTCTAAGCTAGAGTGCTGTGGAGCAATCTTGGCTCACTGCAACCTCTGCCTCTCCTGGGTTCAAGTGGTCTCTGCC  
 7913 TCAGCCTCCCAAGTAGCTGGGATTACAGGCGTGCACCACCACGCCTGGCTAACCTTTGTATTTTGTAGTAGGGACAGGGTTTCAGCCTGTTAG  
 8005 TCAGGCTGGTCTCAGACTCCTGACCTCAAGTGATCTGCCCCCTTGGCCTCCCAAAGTGCTAGGATTACTGGCATGAGCCACTGTGCCTGGC  
 8097 CAGAAAGAGAAATTATTACAATTTAGGTTGTTTGCTTTAGTTTTTCCCCTTGGAGTGTGTTTTTCTCCAGGTAATTTAGGTAGGAAG  
 8189 GAATAATTAGATGTTTTAATTTTGTCTTTAAGTGCACCTTCATTTAAAAATAATGATTTTTTTTTTAATCCTGGTTTTTCTAGTTGATATT  
 8281 TAGATCATAAATATGCTCATCAATAAAATTGCTTACTATAAGGAAGCTATAAATACTCTTATAAAGACCAATTAAATAACAATATTTAATTT  
 8373 CCATTGAGATTTTTGAAAAATTAAATATAAAATTAAAAAATTTTAAGTGTGTTCCCTCATCTTTCTGAAGAAGTAACTTCCTGTCTTACCT  
 8465 CCTTTGCCACTATATTAGTAACTTAATTCAGACAAATACAGCCAGATATGTTTGTGAATGTAGTTATAAATGTCTTTTAAAGGCAGGTAG  
 8557 TGGCAAACTATGACCTGCAAGCAAAATCCAGCCTGTAGCCAATTTTGTAAATAAAGTTTATTGTAAAGCAGCCAAGCACATTTGTTTACA  
 8649 TATTGCCTATGGCTACTGTCACCATGCAACTCAAAGTTAAGTAGAGATAATATGACCTCAAAGCTGAAAATATTTATGATCTTGCCTTTTA  
 8741 CACAAAAGTTTTGTTGACCTATGTTTTAAAGCATGTGGCAAAATTATTAATTGCTAACTCAGTTCTCCAGTTGATTAAAAAATATGGTTT  
 8833 TTTGAGGGAGAACTCTCCATTAAGTTATTTAATCACTGCAGGTTGAGCAATAGCTGCTTCATCCTATGCTGCTGGAGCCAACATAACTAAAC  
 8925 ACTTTTGGGACCCTTCCACTTGGGTGGAGTGAACATCACTTCCTCTTCATCCTCTGATCCAGGGAATGACCTAATGGCTTAAACAAAGCAA  
 9017 AACAAAGCAGAAAAAACTTCAAAAACCTTTCACTGTAACCTTCAAAATATTATTGAATTTCAACAGTTTGAAATGTTAATCGTATAATCAGT  
 9109 CAGTCACATTCCTGTCTTTTTGAAGGTACAGTTCTCAGGATCTGGCTTTCTGATGGAACCTCTTATCTCCTAGACTTTCTGCATCCCCAGAG

FIG. 12B  
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9201 GGGTGGGGGTTGCTTGCCTAATTCTGTGCTTCTGCTTTGGAATTTAGCCAATGCCTGCTATGTAGATGGTCAACACTGGCTTGTAAATCAA  
 9293 TGAATCTCTAAATACTTAGCCAGGTTCACTGTGGAGTTTTTTTTTTGTTTTTTTTTTTTTTGTATGTGCGATTCTTATAATTATAAATATA  
 9385 TGAACATAATAAACTGTAATTATTTTCCCCATTTTGCCTAAGGTATTAAAATCTTGGCCAGGCGCAGTGGCTCACGCCTGTAATCCCAGTAC  
 9477 TTTGGGAGGCCGAGGCGGGTGGATCACGAGGTCAGGAGTTGAGACCAGCCTGACCAACATTGTGAAACCTGTCTCTACTAAAAATACAAA  
 9569 AAGTAGCCAGGCGTCGTGGTGTTCGCCTGTAATCCCAGCTACTCGGGAGGCTGAGGTAGGAGAATCGCTTGAACCCGGGAGGCAGAGGTTGC  
 9661 AGTGAGCCAAGATTGCACCACTGCACTCCAGCCTGGGTAAACAGAGCAAGATTCCTGCTTAAAAAAAATAAATAAATAAATAAATAAATCTGGG  
 9753 CTCCATTCATAAATGCATTTATAATATACAAATTTATGCTGGTTTATTTAGTTTTTAATTGGGCAGGTACATTTTATACTTCAGAATAGTTT  
 9845 TAATGTCTGCAGGAGACTTGGTGGTACAATTTCACTTCTAGAGCTTGTGAAAAACCGCTGTAGCTCTTTGAAGAGAATAGGATCACACCC  
 9937 ACAAGACCACGTCGTACATAGTCAGGCCTGACTACTTAGCTGTGCCAGTGGACCTAGGGGCAGTAGTGGGAAAATCCAAAATATGGATTAT  
 10029 TCTGAAATCTGACTGGTTTTGGAATGAAATTTTGAGTGTGAATGTATGATACTCAAAATGTGAAATGCTTTGAGAAATCTAAAAAATCAT  
 10121 TTTCAAGGATAAAATAGCATTTTAAATATTTCTTAACCTACAGAGTAAAAAATCAAAAAGTTTGCCAGCTCCAAAATCTGACTTTGCGCTTA  
 10213 GCCCTTCTCTGCCATTTATTCACCTCTCAGAATAATTTAAAAACTTACATTACCTTCACATCACACATCACACCTTGCCAGTAAAGTGC  
 10305 TTATTGTAGAGCTTGGCACAAAATAAGCACTTGTGCTGTTGTAATCTTACTTTTCATGAGTTTGGAGGTGGAGGGGAGTTTGCTTCAGAAG  
 10397 ACCTTCAGGCATGGCCTGTGACACAGATAGAAAGGTTGACAGTAGAAACGATGAAGTGAAAGAAGCAGAATAGAGACGTTGTTTCATGTTAGT  
 10489 TACAGCCAGAGCCAAATAAACTAGATAAGGCATAAATACATTTATCGACCTATGTTTTTCATTGATTCACTCATGCTTTTTTCCACCCACAC  
 10581 TGAAGTCTGCACATAAATAGTCAAGTGTGACTGGACAGAAGATCATTTGAACTAAGCTGATTTTAACTAAAATCTTTAGCTTTTGATTTT  
 10673 GACTCATTCATATTTGAAGATCTGGTGGATGTGGCATTATAATCTTAGGCTTTATGCACTATTGGAGGATCTTCATGGGTGGATGAACGACG  
 10765 GACACCAGGCTGGCCCATTTGAATCTGTGATGTGGCGAGGTAGTTTGAATGCAGTCTGTGTAGGTGAGGTCACATAAGAATCCTTGCCAGTG  
 10857 GAAGACTACATTAAACATTTCTGTACACACACACAGATGAAATGTGATGGTGCTAATATTAATTGGAATGCATATGTTTCTTCTAATTATT  
 10949 TCACTCTTTTCTTATGAATGGAAAGAAAACTGAGGGCAGATTTATATGGTAATATTAACAGAGCTAGCTAGTTCTAATATTAGTAA  
 11041 ATACCATGAAGGCGTCTTTAAGTTCACGCATCCTATTCATTTTATGTCATAGTAGAGGCCACATGGCTTTTAAAGAACAGGTGCTTACTC  
 11133 TGAAATAGTCCATTGTTAAGTAAATGTCACAAGGTTAGGTGAAGTTGTCTCCTTGTAACCTGCTCTCAGATCATTAAATGATGATACTTA  
 11225 AAGTGATACTACCCCCAAGGGTAATGTTTCAGTGGTTCAAAGTCTCAAGCTTCACAGGAATCTCTGGTGGTGAATTTAGGCATTAGCTCG  
 11317 TTGATGGGAAAAAATTTTTTGATGTTTCTATGGGTATGCCTTCACAACTTTCCCAAGTGTTTTCCAAAAATCACGTCTTCTGTTTTTTA  
 11409 TTTTCTATTTTTAAATCCCCCTTCAATAGTGCATCAAACCTATAACTCTGACTGGCATCTTGTGAACATAAATATGAAGATTACTCAGG  
 11501 AGAGTTTCGACAGCTTCCGAAGTGAGTAAGCTATATTATACACATAGGGAAAAGTCTTTGTACTTGAAATGCTTGGGGGGAGGTATGTAAT  
 11593 TCATATGCAATCAGAGTAATTGAGGAAAATATTTTAGATGGTTTATGTGTATGTGGTGTAACTATTGTTTACAGGGCCTTTGATTGTAAAG  
 11685 CTCAAACATGCTACTTTGGTATTGATAGGTAGTAATAAATGTTGGATGGTTTAAATTATGTCCAGTGGTTCTTTTACAGGTACCACTGATAA  
 11777 ATAAATAGGTGAAATTTTTTCACTACAAAATATAAAACAAACAAACGTAAGTGTACTGAAATATGCTGCAGTGCATTTTTCTCTTGAAAAG  
 11869 AATATTTTGAAGAAGATAATTATAAAGAGCATTTTACATTGAATAAATTTTATGTTTTTAAAAAAGTAAATCAAGAAACAAGCATTTTCCC  
 11961 AGTTAAATTTTTTTTTTATCTCCTTTAATGTATTAACTTATTCTAGACTATACCAAAGCAAGTGTATTAGATTGAATAGTTGTGGCCAA  
 12053 GTGAATCGCGGTAGCTAGGTATTGCCTTGACAGACTATCTTTTATAAAAGGTTCCATTGTGTGTTGCTTTAAAGGAATACCTGAGACTGGGT  
 12145 AATTTCTAAAGAAAAGAGATTTATTTATTTATTTTATTTATTTAGTTTTGAGACAGAGTCTTGCTCTGTCACCCAGGCTGGAATACAATGA  
 12237 CACGATCTCTGCTCACTGCAACCTCCACCTCCCAAGTTCAAGCAACTCTCCCGCCTCAGCCTCCCGAGTTGCTGGGATTACAGCCACTTGCC  
 12329 ACCACGCCCCGGCTAATTTTGTATTTTGTAGTAGAGACAGGGTTTCAACGTTGGCCAGGCTGGTCTCGAACTCCTGAACTCAGGTGATCCAC  
 12421 CTGCCTCTGCCTCCCAAAGTGCTGGGATTACAAGCATTAGCCACCCGAGACTAGCCAAGAAAAGAGATTTATTTGCTTATAGTTCTGCAGGT  
 12513 TGTACAAGAAGCATAGCACTGGCTTCTGCTCAGCTTCTGTTGAAGACTTTTGTGCTGCTTCAAACATGATGGAAAAGGTCAAAGGGAAAGT  
 12605 TGGCACTTGTGAAAGAGACCAAAGAGGAGGAGGAAACTCACTTTATAACAACCCATTCTCTTGGGTACTAATCCAATTCAGCAACAAGTAA  
 12697 TCCCATCTTGCCAGGAACAAGAAATCACTCATTACTGTGAGAACAGCACCAAGCCTCTCATGATGGATCTGCCCCATAACCCACTAGGCCCA  
 12789 ATTATGCAACAACGGGGACCAAATTTCAATCTGAGTTTTGGTGGGAATAAAAACCATATCCAAACCATAGCAGTTGGCAAATTGAATTATAC  
 12881 TTGATTTTGGGAAAATTGAAAGCAAATAGTGATGGATTATGTTTTAAACTAACCATCACCATGAAATATTACTTGAGACCTGATTAATG  
 12973 TATTTTCTTTGCGTTTGTACATCTTTGAGCTGGAACCTTTTATGCGGTTCTCAGTAGACCTAGCTGTTTGTTTTTCTCCTTGTGTGGCT  
 13065 TTGCCACTCTTAAGAATGTTGGGCAATTTCCCGATTGCCTCTTTTAAACCTCAGCCAGGAACACTCCCTCCTAGTATTATCTTCTCCAG  
 13157 ATGGGTAGCCCTTTAGTTCTATATTTACCCAATCCTCCCTTAGGGATTTTAAATTCTTCCATTGGATTGGCTTAACCATCTTGTGAT  
 13249 CCTCTTGTTCATAGTCTCAGGGTTGAAAGATATCAGACTATGTCATGTCGTATACTTACTATCTAATAGACTGCTGGTACATTTTCTCTCTT  
 13341 GGCATTAATGAGAATTTCCAAATGTGTGATGAGAAAGAGAGGAGAAATTGTAACAGTGGTGAAGAACACAGATTCTGTGTTCTGCCCTCAGAC  
 13433 GACTCTGCCGTCTGTGAGCCTGATTTTCTCATCTGTTAAATGGCCTTAACAACAGTCACCATGATTAAAGGATTAAATGAGAGGGCACATG  
 13525 AGAAGTGCGCAGGGCCTAGCACGTCATACCCATTGAGTAAATGTAAGCTGCTTATTGGTATGGTGTCTTGTTTTTGTGTGGTTAATACCA  
 13617 TCATTGTTAATCGGTTTCAACGCAACAACCTCAATCTTCTTTTTTCTCATAAACTTTGTAATAAAGTTATTCTACCAAGTCTTTGTTTA  
 13709 TTAAAACTAATCCACTTTCTTATTTTAGTATGCCTGCTAACTCCCCAGAAGCTATGCTGTCTTTTCCACATAGCTTTTTGGAGCTTTCTT

FIG. 12B  
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13801 ACTCAAGTCTCTTGGCTTACCCACCTTGAAAAGCAAGGGCATAGATGGTTTTATTCTTTGTCTGAATAAAGAAGCTGGGCCATCTTTGGATT  
 13893 TAGTAAAGGCCGGGCCCCATGATGGAGGAAGAAATGCAAAGCCTCTTCCTTGACTAGGCATTTCTAAAACAAATTTTCTTTGTTGTTTTAT  
 13985 AGCAAAGTGGTCAAGTTGGATAAACTTCCAGTTCATGTCTATGAAGTTGACGAGGAGGTCGACAAAGATGAGGTGGGATACCTGCTTGCTGT  
 14077 TGCTTCTCTTTTCACTCTAGATTTAAACATCAATTTTACAGACTTAGAAGATTAGTTAGAAAATTACCGACATTTAGCCAAAACAGGCATTG  
 14169 GAGTGTACATGAAACGGGAATAATTTTTTAAAAATGTTATTGATTGATTGGAATAAGGTCTCTGTTCAACTTTACTGCTTAGCATTTCAT  
 14261 GTTCTCTTGGTTGTGTTTATTTGTTCTGAGATCATTTTCAAAGACTTGGATCAGATCTGGCTACATTGTTAAAAGATATCAAGATGACTTAG  
 14353 ACCTTGAATTTAGGTTGTTTTTCAACAGATCTCGAAACAGCTGCCAGCCAGTAGATTTAAATGGCTATTTCTTCAATGATTGCTTTTAGTGA  
 14445 AGTCTGATTTGATCAAGCCCACTCCCCCTATTCCTAGAGGAAAGCTCATGGCTAAAGAACTATATAAAGGGAGTAGGGCATTGAGATGAGTC  
 14537 TGCCCACTGAGTGAGGGAAACCTCACAAGAAGACAATGCCCATCTCTGCATTTCTCATCCTCCCCATTGATTGTTAAGTGTCCTATTGTGAG  
 14629 TTTAGGTTTTTCTCTTTTAAAAAAATGTCAGCTGAGCTATAACATTAGCCACTCATTAAGCAATGTGCATGTAGCAAATATTTTTTATTC  
 14721 CCCCCATCACTTTATCTCTCTCTTCTGTATTGCCTCAATTTCTCCCTTGCTTTATTCACCTTTCCCTGAACTAAGCCTCTGGGAAGGTTTC  
 14813 CAGGAATGTGCATGTGCTTTTGTCTCTGACTATAGGGGAGTGTCAATTTGAAAACATTTTTTTCGTGAAACCAGGCAAGACCTTCCAACGTGA  
 14905 GTGGTCAGTTGAGGTATGTCCTTTTGGTCCTTTTGTGGCTCATTAAACACTGACAAATAAAAATTTGGACAGGAGCTAGCTTTGCCTTTAA  
 14997 TGGAATAAAGTTTTTCAGAAATGTAGGCGGGTCTCTCTCTTTCCACCGCTAAGTGGACTTTTTATGTGACTTGTAGGCATTGGTGTCAATGGGTG  
 15089 CTTCACTAAAGGGCAATGGACAACCTGGCACAAGGGAATGACCTTCCCATTGACCAAACTCACAGCAAGCAACCCAGGTAATAACGGGAG  
 15181 GTTGACTGCTGGTGTTTTCTCTCTCTAGGATGCTGCCTCCCTTGGTTCCAGAAAGGGTGGGATCACCAAGCATGGCTGGCTGTACAAAG  
 15273 GCAACATGAACAGTGCCATCAGCGTGACCATGAGGGTGAGGACGCACATCACTTTGCCCTCCCCTCTCACAAGCCCTTTCTGCCATAGAGCT  
 15365 CGAGAACAAATGCTCAAGATGAATGCGCATGCTGTTCTTCCCCACAAAAGGGACATTGTCTGATTCTTAGGATGCTCCCCTGGTGATAGCACC  
 15457 CCCATTGGCACAGCCTCATCCACCCACTTTCCCTCACTGTCTTCTGACCACCAGCATAAGGAGACCATCCCTGGGCTGGTGTGAAGGTGCAG  
 15549 AACTGACATAGGCTTTCTTCTCTGTAATAACTGAAAAGTGCTCTTTGGTACCTCACAGAATGTCACCAAGGGGCTATCTGTCTATGCCAATC  
 15641 CTGAGCACTTCTGTGGAGGTGTACTGCAGCAAAGTCAAGTAAAGCAAAAATTGAGGACGAGAAAAGAAAATAGTTGCATAGAAGAGAAGGTT  
 15733 GCAGACAGAGAAGTCAAACCAATAGAAGAAGCTATTCAAGGAGAAAAGTGGGACCAGAGGAACATCAGGATTAATAACAAAGGGAAGAGAAAC  
 15825 AAGGGAGTCAGGGAGATAAAAATTAAGGAGGAAATGTGACTGTCATTACCCTAAGGCTGGAAAATCATTGAGCGTCATGAGGCAAAAATAG  
 15917 TTCCCATCTGTGAGCAAGAAACCCTGGGGATTTTAGAGAAAAGTTTCTGTCTTCTGTGCTGCATCCCAAATTGGAAGTCCCTGCACTGCTT  
 16009 TTGGGTAGTTATGTAATACTCTGATTCCGTGGGTGAGAAAAATGACCCATGGATATTAGGGGAACCACCTCCTCAGAACTGAGATGCAGTG  
 16101 AGCTTCTTAGATGGGATGGGGAGTCTTGACCCACAGTGACCTGGAGCATCAGCTAGAGTGAGAACGGAAACAGGTTTTATGTATGTATGTA  
 16193 GTCATAAGTGGGTTATTGATAGAGATTGTGACCCCTCTTCATTTTGAAAGAATAGCTGTGTGTATATTTTTCTCTCAGTCATTTAAGAGACGA  
 16285 TTTTTCACCTGATTCAACTTGGCGATGGATCCTATAATTTGAATTTTTATAAAGATGAAAAGATCTCCAAAGAACCAAAAGGATCAATATT  
 16377 TCTGGATTCTGTATGGGTGTCGTTTCAAGTAAATATGAAAAGAGTTTTACCATTATGTTTTCTTATCTGCAGTAGTGCTTATGTGTAAATTA  
 16469 GCAGATTTAAGCAAACACTTCCAAAATGGCAATATGCATGGTAGAAATATAACATATAACTTTAAATGAGGCAAGCCTTGTTTTTTCATCAT  
 16561 TGTAGAAGATGGAAGGGATAATGTAGAGGCAGAAATATGCTGTGGCAGGCAGGAGCACTCTGGCTCGGCCACTTTATAGCTGCGTGACCTTT  
 16653 AACAGGCTACTTAATTCAGATAATGAGAATGTTTCTTTAATACGGCAAATGAGTACATTGGATGAATCAGTGCAGGAAAATATTTAAACAC  
 16745 TTCATAGTATCTCAGTGGTGATTTTTATCGCTAGCATTGTAGTACCAGTGGCGGTGTAGATCAGTAAAGAGATTAGGTTTCAGCGCAGATTG  
 16837 AGTTCAAATCCCTGCTCCTCCACTTACCAACTGTGTAACCTTGGAGATGTTATTTAACCTCTCTGTACCTCAGTTTCTTCATTTGTTAAATA  
 16929 AGGATAATGGCAGTACCAAATATGGTTACTGAGAGGGTTCATTTCATTACACATGTAAAAAGCTTAGAACAGTGCCAACAAATGGTAAGCATT  
 17021 TGGTCAGTATTAGATAGTTTTGTTATCATAGGGCTGTTGTACTTTTATATCATAGGGCTTATGTACTTATCCTTTAAAATTATTGTTAATTA  
 17113 AAGATAACACATGAATGTATTTTTCTTGTAATAAATCAGCCAATACAGATAAAGTGAAAGTCCTTCTGGACTCCTCCCCCTCCTCAGTGTCT  
 17205 CTTTTCTGAGGGGAGCTACTACCAGTTTTGCATGCATCCTTCTGTAGCTTTTTAGCATTTGCTTTTGAAGAGAGTTGTCAATTTCCCTGTCC  
 17297 ATCATCTGTCCATCCATCCATCCATCCATCCTGTCCACCCCTCCATTTCATCCAGCCTTGCCACTTTCAAGGAAGATTTAAGGCAGCAGC  
 17389 TTATAAGCATACACAGGACATGGGATAGCATAAATTTAAAGTGGGGGTGAAAGCAGAAAGATGAACAGGGGATTGGGATAAGGGTGAGAGAA  
 17481 AATAGAGTTAAGGAGAAAGCGTATGTTTTGAAGATCTAACACCTGCTGTGGGTGGGCCACCACCTGGGCTCTATGCTTTCTCACTTGGAGAC  
 17573 CTGTTTAGTCACGCAATTCACAGTGCACATGAGATAAAGGCATGATGCCTGTTTAGTCGACTCTAGAAGCACCCCTGACTTTAAAAGAAGT  
 17665 TAAAGCAAACTAAATGTATTTGGCAACCTCATTTTTTAAAGTAGGAAGTAATTATTTTTGTTTTATAAGAGAGTTTTGCTGCCTGTTTCTG  
 17757 GCCCAGGGACAGATGTTTATAAGTACAACCTGCCCTGAGCTATCAATTAGTCTCCGGGGTGCAATTTCAAAATCTAAGGTTCTGACTTCAATGG  
 17849 AAGTCTCTTCCTTCAAATGTCTTTGCAGATGCAGCTGATGGTGTTCATTTAATAAAGTGTATCCAAGGCTTCAAAAAGTAAATAATT  
 17941 TGTTTTTATCTGTGTCTGTTTGTAACTAAGCATCAAAGTTGTGATTTAAGTGTTTTTTAAAAATTATTACTTATGGATATTATAAAAAAT  
 18033 TAGTTGACTGGTGCTGTGAATTAAAAAAAGTGCCTAAACTAAAAAATTTTGAAGCATTTTAGAACCTTGAAATTTATTATACTTATTTTGC  
 18125 AGATGAGAAAAGTGGGCTCAGAAACAGAAATTTAGAATTGAGGCCAATGTTTTTTCTCCACTTTTAACTTTCTCTTTTCATGATTGTGA  
 18217 GTATGCAGGGAAAGGAGGAGAGAAAATTCATTTTGTTCAGCCTTTGACTTCTTCCCTGGTCTTGCCTTGAAGTTTAAAGTGAATCCAAA  
 18309 GTGGCAATTACTGAGCCACAGCAGACAGTCTGTGCACAAGAGTGTGTGGCTTTGCCAAGGGGAGCACTTGACTTTGCATTTCTAAGAAGTGA

FIG. 12B  
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18401 TGCTGCAGAATCACAGAGACTTTTGGGAGGGTTGCCCTGTCCCTGAGACCTCCACCAAGGAACTCTTAGAGAGAGTGTGGATAACCCAGTAG  
 18493 GATTTTAGTGGCTATGCGGGGGGCTGTCCTGCTGGCTCAGGTTAGTGGGAGTGTGTTGATTTTCATATCGCTCAGCCTGTCCTTACAGGGGATC  
 18585 TTGTGCCATGATCCTCAGAGCTGAACCTCTGTCTACTGCGGCCAACCTGGGGAGATTTTGCTCCCTGGAGGACATCTTGAATGTCTGAAGA  
 18677 CTGGCATCTATTGGCTTAAGGCCATAAATTTGCTTAAACATTGTACAATGCATGGACCAGCCACTCACAACAAAGAATTGGCTGCCCAAGTG  
 18769 TCAGTAGTACCGAGATTGAGAAATCCTGGCCTAGTGCATGTTTCATCTTCCGTCTGTTACTGCACATGGACTACTGTTCTTGTCTGTGAGCC  
 18861 AGTCACCTCTTGCAGGCATGAAACTGGAGGCATGAGGCAAGGCCACGGACAGGGAGTCCAAATACCTTTTGGGATTCATAAAGGATGGGA  
 18953 AAGTTCAGATAAGTAAGCCAAACATAGTAATAGATAATGGTTGGCTTTTAAAAATGTAATACCATACTACTTCATTAAAAAATAGGAG  
 19045 CTGAAGAAATATGAAATTTTACATGAAATTTTATTATTCAACAAATATTTTCAAATACCCACTATGTGCAAGTCACTGTAGAGTCCATA  
 19137 GAGACTAAGGATGTGTAGCACTGACAAAAATGGGAGCACTGAGGAGGTTTCATTCCACTGCAGGGACACACAGTGAATCAGATGAGTATGTA  
 19229 AAGCAGGTAATGAGTCAGAAGGAAAAATAAAGCTTGCAGAAAGTGAAGCAGGGAAGGTGGACGGTAATGGGATTTTCATGGGGGGGGCTTTC  
 19321 ATGAGGAGGGGGCAAGCTATTTAAATAGCTTGGTTCTAAATGCCAATGAGATATCACTCACCACAAAGAGAGAGTAATTATTTTAAAGCAG  
 19413 TTCTAATTCTTTTAAAGTATGCTGTTTATCCTTTTTCATTTTCAAGAACAAAGTCAAGCGTTTGTCTTTGAGCTCAAGATGCAGGACA  
 19505 AAAGTAGTTATCTCTTGGCAGCAGACAGTGAAGTGGAAATGGAAGAATGGATCACAATTCTAAATAAGATCCTCCAGCTCAACTTTGAAGCT  
 19597 GCAATGCAAGAAAAGCGAATGGCGACTCTCACGAAGGTAGATAGGCTTGGCTTCCCCAGGCACATACACACTCTGTGGGTGTCTTTATTT  
 19689 TTGCCAGGTGGGTATAAGAAGGAGACCTGTGTTACACAAGTACATGAGAGGTGGGACGGATAGGAGCTCTTACAAATATCCTGTCAGCAAA  
 19781 GGTTTTGTACATTATAACTTACTTCCCTGACATTTCTGTATATGGAATCATGTAATGGGAAGAACCAGCTTTGGAGGCAGAAAGGGAGA  
 19873 CCTGGGTTTGTAGTCCATAAATACTGTATTTTCACTGTGTAGCCCTGGGTAAACAACCTTATGTTTTCTGAGCCTCAGTTGACTCACCTATAA  
 19965 AATGGGAATAACATGAAATTTGCTGGGAAGATGGGAAGTGTAAATAAGAAAATGAATCTCAAGTATCTGGCATAGAATTTTACTGTATTAT  
 20057 AAAATATTAGTAATAATTAGAATGCATGGGAGCCTCAGATTAAATTTGGTGAGAAAAATCTGGCTATGTTCTTGACAATTCATGTTTTACTTC  
 20149 AACCTTAGGTGATTCCCAACCTGGCTTCCCTTAGAAGTACCTGGGAGCTTTTAAAAATACCATTTACCTGGTCCCAAAAAGATTCTG  
 20241 ATTTAGTTGGTCTGGGGTGGAGCCTGGGCAGGTCTGACTTTTAGGGGGTCTCATGGACGTGTCCATGTGGGCTGTTGTTTCATAGCTAGTGTC  
 20333 AGTTCTAATTGGACGGTGTCCATGCTATACCAGCTGCTCAGTGTGTTTACTTTTCATCACTGAGCCTGTGGATCAGTATTTTTTCAAAGCACC  
 20425 CCAAGTGTGTTCCAGGAGCATCCAGAGTGGGAACCACTGTGTTTCATTTGAAGGCACCTAAGAGAAACGGCCTTCTCCTCCTGTTTCAAT  
 20517 GAAATGCTATGAATTACAAGTGATTCCGATAATCTGTTTTGCCATTTTAGATGATGAACAAAGCAAATTGGAAGGTTCTGGTTCCGGTTTAG  
 20609 ATAGCTACCTGCCGAACCTTGCCAAGGTAACATCGTCTTATATCTTCTGCTCTTCGTTGAATGCTGTTGAAGTATGTCTCATTTCACTGGTT  
 20701 TGTCCAGAATGGAATCTGTTGAAATCATAAAAATTACATTGTGATTACCTCTCTCTTTTTCTGACCTGATTACGAGGTGACGTGTACTCATG  
 20793 CAGTATGATTTTCAAGTCTGTCTTCTAAAAAGTACCTTACAAAGCATTCCTCTTTTATTATTATTTTAAAGTGTTTTTTCCCTGATAATGCTT  
 20885 AACACTGCATCACAGGTAAGTGAAGAAATAACTGAAATATGCAGGCAGATGTTCTCATAATAGCATCGTACTTTCTATGTTGATACATGTGCT  
 20977 CTCCCTTACTCAGGGTAATAGACACGGTTCCAAAGAGGAAGGACCTGGTAATCTTGCCACGAAACCCGGGGGTGCTGAGTTACAGAAATT  
 21069 GTTTCGGGTCACCTTACTGGAATAAATAAGCTATTCCTGTGTCTTACAATTTTGAAGAAATTTAAAGTTACTGAAAAGCACAAAGAAAA  
 21161 GCAAATCAACCATACTGCTACTTCCAGATTAAATATCTATTATGATGTTGCCTTTTTAGCTTCCATATTCTTAAAGATATAAAACATCGG  
 21253 TTATAGTTGAAGTTCTTTTTTAAACACTGTCCTTATTCCCATCTCTTCTCTCCCAACCCCAATCAGAAACAAGCACTATTAAAGTTTT  
 21345 ACTTTTCATTTTATATCTTTACAAATAAATCTATCATAATGATATATACAGTATGATTTAGTATGTTTAAAAATGTTTTATAAATGCTAACA  
 21437 TACCATATGTATTCTGCACTTTAAAAAATTTTTAAATTTACCCCTTTTATTTGTACTATATAGACTTTTTATTTTAGCTGTTCTATTATTTT  
 21529 CATTTTTTCTATTATAAACAAGCTACAATGACTGTCCTTGTACTTGTGTCCTTGTGTGATCTGAATGATTCTTTCTTAAATGAGAGAAA  
 21621 TATCTTTGTCTTCTCTAGCCCTTTGCACTCTTACTCTGTTACTGCCCTTCTATTCTTTTTTGATACTAGAGTGAAATGGCGACCCTCCACAC  
 21713 CCACATCTTAAACACTATAATAGAAACATGGTTTATCTATATAGGATTATAAATAGACCAGCATTGAGCATTGACCTTTATTTTAAGACAAC  
 21805 ATGGCTGTTCTCAAGTGTAATCTCCCTCCCTGGCTAGGGCTTTAGAGCATTGTTTTTCTTTAGGACTTGACTGCTACCACAGTATCTTTT  
 21897 TAGCACCTGCCTATTAAAGCTAATTTTAGTGCCACCATGTAACCACCTCCTAGTCTGGGAAGAGTTTTGGCTTGTGTGTTTGTGTTATGA  
 21989 ATGTCTGTGTATCATATTTTGCAATTGAGATTTGCTTTTTTGTGTTCTGGATGTTTGGGGGTTTCAATTTCTCAAAACAAATATTTGTGCCC  
 22081 ATTTGGGTTTTAGTTTGTGTCAGCAGGTAATATATGTGATGCCATCTAGAATTCAGAAAGTAACCTTCTGCACTTACTGGGTGAACGGAATG  
 22173 GATACCTAGGAGAAGATTCATGTTATTTGAGCCTAATGTTGATTAAATAAAAATCTATGCTTTTCCCTATGAGGATATACAGGAACGGTCCC  
 22265 CTCCTTCTCTTACTACCCAGCCAATATAATTGAGTATGTTTGTATCCCAAGACCTAGGGAGATTTTTTAAAGATATACATATATTTAATATAA  
 22357 ATGTATACATTTATGTATATATACTTTTTTATAAGTATAATTGTATATTTTGTCAATTTAAATATTTTGAAGTATTTTTAAACTATGGGTAA  
 22449 CAAGTTAGGTTAAGCCATTTTAGTTGGTGAAATCAGTTGATTTCAACCCCTGCTTCTTTTTGTTTTGTTTCTATTAGTTTTTTATCTTT  
 22541 TTATTGAGGTATAATTTGCAATAGCAGAAATGCTCAAAACATGAATTGTAGAGCTCACTGGAGTTCGCAATTTGTACACTGATATAAGCAGCCCT  
 22633 CAGGCTAGCTTGTACCTGAGACCCCTCTTTATTTTGACCTCCATCACCGTAGATTAGTTTTGACTTTTCTAGACCTTCTGTGAATGGACTTA  
 22725 TACATGTACTCTTTGTGTGAGGCTTATTTAGCTAAACATGTGATTCATTTAAGAAGTTTTTTTTAGGTCGGGCATAGTGGCTCATGCCTGT  
 22817 AATCCCAGCACTTTGGGAGGCTGAGGTGAGCGGATCTTTTGGGTTAGGAGTTCAAGACCAGCCTTGCCAACATGGTATAAAACCTGTCTC  
 22909 TACTAGAAATACAAAATTAGCTAGGCGTGGTGGCAGGTGCCTGTAATCCCAGCTACTTGGGAGGCTGAGGCAGGAGAATCATTTGAACCTG

FIG. 12B  
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23001 GAAGGCAGAGATTGCAGTGAGCTGAGATCATGCCACTGCACTCTAGCCTGGGTGACAGAGCAAGACTGTGTCTCAAAAAAAAAAAAAAGGGTC  
 23093 CGTTTTAATGAAATAAAATGGAATGGAGAATATGAAAGTACACTGCCCTTAATAATGACATTATTTTTTATATAAAATACTGTCATTATTAT  
 23185 TTTGGTGGCACCTGCCACCATGCCCTAGCTAATTTTTGTATTTCTAGTAGAGACAGGGTTTTATACCATGTTGGCGAGGCTGGTCTTGAACCTC  
 23277 CTAACCTCAAAGATCCACCCACCTCAGCCTCCCAAAGTGCTGAGATTACAGGCATGAGCCACTACGCCCGACCTGAAAAAAAAACTTTTTAA  
 23369 AGTGAATTACATAATTTTTTACATAAAATAATGTCATTATTAAGGGCAATGTACTATTTATACATATAGTGTGTATGTGTGTCTTGCATAGT  
 23461 GATATAAAAGATATTTGTTTTCTTAGTGTGCTATTATGTATATTTATTTACTTTTCATTGGTATATAATGTACCTATTTTGGGAGTTCATGT  
 23553 GATACTTTGATATCTGTATACAATGTGTGATGATCAAATCAGGATAATTGGGATAGCCATCACCTCAAACATTTATCTTTGTGTTGGGAATT  
 23645 TGAAACATTTCTTACCAGGAGTCATGGTCAAACCTGAAAAATGAATCCTTGTTAGAGGCTTTTACTCTTTCCCCCTGGCTTTTCAGGTGTTT  
 23737 TACAAATACTTTTATTTAGGAAGGTAGAAAGGTGAAAGTAATTTTTTTGAAGGGGAAAAGAATGAAGAAAATGGAGATGAGTTATTCACCTCA  
 23829 GCACATGGGTATCTGTGGGCTTTGCCTTTTAAAGCCCAGCTTGGTGTCACTGTGAGCAGCCAGGCAGTAAGGGGAGACCTGTGTTCCCCAT  
 23921 CCCCAGCCTTGAGCAAAAATGCAGTTTGGCTGTTTATCATCCCCCTTCAGGGTGTCTGAACTATTTGCACCGGTTGAGAAGGCAAGAAGT  
 24013 TGACCTGATAACTGTTGGTCATCCCATTAGGAAGGATGGATTCCATGGTTACAGAATCAGAGACTGAAGTATGCAGAGGGAGGGGTGGGGAG  
 24105 AGAGAACTGTGCAAGGAGTTTACCCAGGGTATGAAGAGGTAAAGAGGTCACTATCAGGGAAGGAAGGTGCAAGAAAGGTCAGGCTGGGAGG  
 24197 CTGGGCCACAGTTCAGTAAGATTACAAAGAAGGGCCTAGAACAAATGAGGGCAGGCAGAAGGTGGCTGAAGGTGTAATTTTCATGGCAGGTTCC  
 24289 TTTTCTAATCAGCTCCTCTAACCTCCTTCATCCTGTTGCCCGGGCTTTTGTTTTCCACTGTGACTAAGACATAGCCAAACAGGATATGACCG  
 24381 ACAGGAAGTTGTTTCAGTGCAAAAATAACTGATGTCTCATTCTGGAATATATGGAAGGGCTCATTACTTACAGTGTGAGTGATGTAAACCC  
 24473 AGGTTTTTCAGAATATTTTGTATAATCTTGGAGCTTATGTTTGTACATTTAGTACTGAACATCTGTATTGTTTTCTTATTAGAGAACACACTG  
 24565 TATTTACCCATAAACTGGTTCCTTTTCCTCCTATTGTCTATTATGGAACCAACAATTTTTATTGTAAATGTAACAGTGTGTAGCATCAGTCT  
 24657 TATAAATATTTTAGTTTGTATACACAAACCGTAGTTCAAGTTAGTTAATTGATTTCTTCCCTAGAAAGTCAAGGAGTAACATAATCAGGTTAT  
 24749 AAACCTCATTACTAGTTATTTAATAATTTATTTCTCTGGTTACATTTATATCTTAGGTGACATCAGAACATATATGTCACCTCCTTAAAGAT  
 24841 AGTGTGAAGAAAACCCACCTTATGTTTTCTTCCACAGCTTTTCTGTTTGTGAGCTTTTATTTTGTACTCAAAGAATAGCATCCAACTTTTA  
 24933 CTTTGGTTTTCCCCATGTGGTTCTGAAAGAGAAGTAGAATTTCTTCTAAATCCGGAATTGCTCACATCCTTTTACCTTTTAACTTTGTTTTAA  
 25025 GCAAATGAACCTTATTGTTCCAGGTAAATCTTCCACAGTTGCATGCAGGGGAAAGTATGATGTCTCAGACTTTATAGTCTCATGGAGATGGAG  
 25117 TGAGGATCAAGGGCCATGCTCAGCAGAACTTGTGAGGACCCAGCAGTTTACGGACACCTTTTCTTAATTTTTTAAACCAAGTCTATAATAAG  
 25209 TGCTTTCTTCCCTAGATTTCAATCCAGAAACAATATCATTGCATATTATACAAAGGAGCTGGCTAGGCTTGTGTCTGTGGGGTCAGCTGG  
 25301 TGTTGCATTTCTGGGCCTCCTTTGTGAAGAGGATGAACCTGATGGTCTGAGAAGTTAGGTGTCTTGAAGTAGTGGAAATAAATCATGATAA  
 25393 CTCTTTAAATTAAAGATTATATATTTTGGCCTCAAAACATTTTGCAAAGTCTCCTATTCCAACCCAATCTGTTTAAATGACCCAACATTCA  
 25485 ACACATTGTTTCTGATAATTCATCCTCAGAATAAGATGCTGTTGGCCATAATCTTTGTCTCTAGATTGTTTTATCTACTCGAAATAAATTT  
 25577 AAGACACAGAGTATGCCTAAAGCCTACAGCAGACTTTCTGGAAACTCTTGAATGTTTGGTCCATAACTACTTCTTAAGACAAAGAAGAAAAC  
 25669 CTTGTGAGGGTGTGTCAATTAGTGCTTGAATGTAGGGTTTACAGGATGGGGTGGGGTGGGGGAATCGCCCTTGGTTTAGATGAATCATTTCTT  
 25761 TCCTTGTCTTCTCAGCAAAACACCAGTTTCTACAGAGAACAGCTCTGCCATTGTGCATTTTCTGTCTCCATTTTCTCTCATTCTCCTCTCCA  
 25853 CGAAACCCAGAGTAGTCAGTGGGCTTTGGGCAGGAAAGTGGCAACAGGGTGTCTGGGGAAAAGCCAGTTGGCTCTTCTTACCATCACAAATAT  
 25945 AGACTGACCACAGGTTATTTTAAAGAGCAGAGCTGGTTTCCATCACTCTGAGAAGTGCTCAACTACAGACTTTGGGATGATATTTGTTATAGC  
 26037 TGTATTTTCTCCACTCTTAGATTGTGAAAGTACATATTACAAGTATTTATTTTATTATCTTTACTAAAATTTTAAATTAAAAAGAAGCGTGCT  
 26129 TGCCGCAATAAGTAAAAATACCCAAAGTTGTTTAAAGAAAAGTTCACCTTTTCCCTTCATCCTCCATTCCACATTCCTGAGAACACTGAAG  
 26221 TTAATAACCGGTTGCAATTCCCCCTTTCACCAAAGTATTGCTCATAGAAATATAGATAAACATATGTAAGGTTTTTAAAGTTTTTTTAAATAAA  
 26313 AATATGTTTCATGATATATACATTATCTGAAATTTTCTGTATCACTTAAAAATAATTCATAGATGTCCCTCTGGGCCAGTGGAAGATCTGGT  
 26405 TCCCCCTTACATACATATCAGCAAGCTGCATGATATTTCAACTATTGCTACTGCACAGTTTATTTCAGCCATCTCCCTATTAATGAACATTTA  
 26497 AGGTTTTTTTTTTCAGTTTTTAGCCTCTACAAACAGTACACAATAAACAACATGACATTAAATACTTGTGCTCTTATTTAGTAGGAGAAATT  
 26589 CCCCATGTGGAATTTTTAAGTCAAAGTTTATTGTGTTTTTAATGCTTTAAACATTGCCAGGTTACCGTCCCAAAAGGCTATAACAATTCAC  
 26681 ATTTCTGTTTCTCTGCATCTTACCAGACGAGTGTAATAAGGTATTTTACTGTGCTTTTCATTTATATTTTGTGTTATTAGTGATATTTT  
 26773 CATATTTTCATATATTTATTTGCCATTTGTGTTTTTCTTCCCTGACTTGCTTGTTCACATTGTTTACCTTGTTTTCTTCTGTCTTGTGTAGT  
 26865 GTAATAGTTTAGACTCTGAAGCCAGGCAACCTGAGTTAGAAGCCAGGCCTCTATTTTCATGATGTAGGTCTTTGGGCAAAGTACCTAACATTC  
 26957 ATGCCCTTAGTGTTTTCTCTTTTAAATGAGCAGGATAATAATAGTACCTGCCCTCCTAAGGTTGTATAAAATTAAATGGGCACCTAGGGTAAT  
 27049 ATCTAGCAGGTAGATATTGGCTATTATCAATAGTAGCTCTTATCGTTACTATTTCTCCAGATACTGTTTCTGACTCTGGGGCAAAGTCCTG  
 27141 CTACCCCTGAACCACATTTTTCTACCTCTTAGATTTTACTTGGTAATTCATCAGCCACTGTTGGGCATCCTCTGTGTTTAAATGCATCATCT  
 27233 TAGACCTTAGGAGGGATGGGAGGAACTTTAAAGAAGCCGAATTTGCTTTTTATTTATCTTGTAGCAGAGCAATAGATGTATATTAGGTAGATT  
 27325 ACAAGCTTTTAGGTTATTTTTGCATCTAAAGCTGTCCCTTCTTTTCCAATAAATGATGTCTGTGGTAAAGAATATATCTGTTGGGTGTTAGT  
 27417 GACAAAATCAGAAATGCTTTGTGTCTATTTTGGCTAGTAGTTAATTGTTTTCTTTTATTGTGTCTGCATTCTATTTGTTCTTTAATTATAC  
 27509 CGAGCTCATTAGCAGTTATTCTTGCTTTATTCATTTCTTATCTCCTAGCATAGTCAGCTCAAGACAACAAGCATCTTTCAGAAAGCCACTAG

FIG. 12B  
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27601 GAATTGCATCTACATTAAGAACCCACTCCTCTGCTCTAGCGTCTGAGAACATAACACAGTATTTGCCTTTGTTGAAGGGCTCAAGCGGACGA  
 27693 TTCAGAAGTAGAATAGATAACTGTTGGTGGTGGCTTGCAGCACCTAGTACTGATGGTTTTGTTAGGAAACACCACAGGCAGTATACAATGTA  
 27785 AAGCAAGTTCTTTGGGTTTCAGTTAAATAGATTCCACTTGCACGTGTTCTCACTCTTTTGGTGTGAAAAATTAGGAAAGGTGATAGGCAGGAT  
 27877 AGAATAAAATGAAGTGGTCCCCTCTCCCCTATGAGAGCGCCTGCCCTCCACCACAGACACATGTTTTGCCCTGGAAGCATAAACAGAAGATT  
 27969 GCAGGAACGCCCCCTTCACCTCCATAGCCTTCAGGCTCCCCTCGATAGCATCAAGATAAACTTGGTGTGGCAACAGACTTGAGCCATCATCTT  
 28061 GTTTAACATTTTTACCTGGAAGTGAAAATGGAATCCAGAGAGGCTAAGTAGCTTGCACAGCTACTTAATTGAACTAGAACCAGAACCAGTT  
 28153 TCTTAGTATTCTGGTGGCCATTTTATATAAACATAGACAGCTGATCATGGTAGTTGGATCATTGCTAAAGACCTATTATATACAACAATCG  
 28245 TATGGCTAAGAAGATTAAGATGCTTATTTTCTCGTTATCCATGATTAGAAATGTAATAAGAGACTTTGTTCTGTCACCAACAGAAGAGACAA  
 28337 GACGCCCTTTCTGCTGTGGTTGTGAGTGCCCCCATCAAGTGTGAGATGTGACTCATGTCTTTGGGGAAAGTAGCAGATGAGAGAGTACATTC  
 28429 TATTCGACTCCATAGACTTAACTGTGGAGACTGAGAGTAGTGAGAGCTCAGACTGACAAGAAGAACAGAATATAGACTTAGAGGCACCAG  
 28521 GCATGAAAAATCATAAAGATGGAGATGACTTCTATCTTGTGAATGTCTGAAAAGCCTCTCTCTCTTTCTGTTCCCAATCCTTCATCTCCAG  
 28613 CCCTCATCTCTGCCACCATCTGTGCGTTTTCTTCCGTGGCCTCTCTTCTGTCCACCTCTTCAATGTTGGGCCCCCAGGATTTTGTCTTCAGA  
 28705 CTCACCTTTCACCTCTCCCTCAGGACATCTCATCCACTCCCAAGGCTGGAACCTCCAGCCTTCCTCTCAGCCTTGCGCTTAAGTTTCATCCCT  
 28797 TTAGCTGTCTATTGGATGTTTCCAGTTGGAAATCCACAGGCATCTCTACCCAGGCGTCTCTACACAAAGATGATAAAAAATATTATCATCTT  
 28889 TTTGGCAGATTTGTTTTCTGAGTTTTCTGTCGTGTTTCATGGATTACCATTCAGCAGGCTGCCCAAGCTAGAAATGTGGGATTTGTTAAT  
 28981 CGGCCCTGCTGTGAGACTGGGAGGCTGGTGTGATAACCCAAGAAAGACATTGTGGGCCTCTGGGTCTTATCCTTTTTTAACCCATTGCCTCAG  
 29073 CCCTGCCCGAGTGATGCTTCTGAAATGTGGAACCTATTTTATTCTCTTACCTAAAAGGTTGCAGGATATTTTAGATTCTGAATGAAATCCCC  
 29165 AATCCTTTTTTTTTTTTTTTAAGTGAAAGCAAGTTTATTAAGAAAGTAAAGAAATAAAGAATGGCTATGCCATTGGCAAAGCAGCCCTGTGG  
 29257 GCTGCTGGTTGCCCATTTTTATGTTTTTTCTTGATGATATGCTAAACAAGGGGTGGGAAATCCCCAATCCTTGGCATTCAAATCCCAGACT  
 29349 CATTTCTTTTCACTTTTTTTTTTTAATCATGCCCTGCCTTTCAGTTATGATGAGTGACTTGGTCATTGCTCAGGTGTGACTTGTCCCTTTGC  
 29441 ACCTGCTGTGCCTTCTCCTAGAATGCGCTTTTCTCCTTTCTGGCCAAGTGTTCTTGTCTTCCAAATGGGCCTTCCCTTGGGAGGTGGTTT  
 29533 CTGACGACCACCCCTAGTCCAAGTCAGCTCCCACTGTACTTTAACTTTCTCTTGTCTTCCCTTATTACCTGTTGATATGCTCTCTCCCCAC  
 29625 CTGGTGTTCCTTGGGACTAGGGACTTCCCTCATTACATTTACATAACTTGAGGGCCTGGCTCATAAGAGGTGCTTAATGAATATTTATTG  
 29717 AATTAATTAGCATCTTGTCTTCAAGATCAGCCATCATTTTCTCTATCTCATCATTCAAATATATTCCCTTCCCTTCTTCCCTTCTTGACCCC  
 29809 AGTCACAGACTGGACTCTATTAAATCCTGTCTATCATCTGGGCTCATTTCCATCCTCAGTGTCTGTCTGTGCATCCTTTTCATTACGCCAGG  
 29901 GATGTTACAGCTCGATTCTGCCCCCTTCATTCCAAGCCTGTCCCATATTCCATTACTTTATGAAGCCTTTCTTGACACACAGATGCTTAATTAT  
 29993 TCTCTTTTGCTTTCTTTGTGTTGACTTTGACTCTGCCACTGGTTGTGAGCTTCAGAAGGGCAGGGATCTCACCTTCACCTTCTTTTTCTCCT  
 30085 AGTGCTTTCTTTGTGTGCTGCACACTCCCTGGCACACACAGCGGCTCTCCAACACGAGGCAGAGCTTTCCAGCAGCCTCAACCTTCAGGACT  
 30177 GGGCAGCTTTTAAATGTATTTGGGCACCTTTGCAAGAAAAGGATTGTGTTAAATGTAATTTTCATGTATCTTGTATCAGAGTGCAAGAGAA  
 30269 GCAGAAATCAAACCTGAAAAGTGAAAGCAGAGTCAAACCTTTTTTATTTGGACCCAGATGCCCAGGTAAGAACTATCTAAATGTTAATATTTA  
 30361 AAACCAAATGTGGGAGAGAAAATCATCGATGGGCTTATTTGTTTATTTGTTTGCTTTGTTTATTTTGGA AAAACAAGCAAATAACTATAGA  
 30453 AAATTTGGGGAAAAGAGGAAAAATAAAATGTATAATCTTATCACCATAGCATTACTATTGTGAATATTTGATATTCAATAGATGTTTGAAA  
 30545 ATTGGGAGAGATTTATTGAAAGACATTCTCAAGTTCACAAAGAACATCTAATTTACCTGTTAAAATAACCATCAGAAAACAACAGGTATCAC  
 30637 TGCAGTTGCCTGGGAGTCAGTGATAATCCCGACTAGCCAGGCTCAGGCTCAAATACAAACCTTTTCCATTAACTCTAACGATAAGTACT  
 30729 TTTCTGTTTCTCACAACCTCATAACCATAACGTATGTGTGTTTATATGTCTATATTTTTTATTTGCTTTTAAAGAGTTTTTGTATTATCAT  
 30821 GTAAATATACATAATATAAAATTTACCATTTTAACCATTTTTAAGTGTACGGTTCAGTGGCATTAATAACATTCTCATTGTTGTACAACCA  
 30913 TTACCACCATCCATTTCCAGAACTTCTTCATTTTCCACACGGAACTTTGTATCAAATGATAACCTTCCCTTCCCTTCTTCCCCCATCCCCCT  
 31005 AGTAACCTCTGTTCTACTCTGTGAACCTGCCTATTTTAGGAACCTCATAAATGTGGAATCATACAGTATTTGTCCTTTGTTTCTGGCTTCTT  
 31097 AAACCTAACATGTTTTCAAGGTCAATCCATGTTGTAGCATGTGTCAGAATTTCCCTTCCCTTCTGTGGCTGAATATTCCATTGTATGTATATA  
 31189 CTACATTTTATATATCCTTGTAATCTGTTGATGGACACTTGGTTGGATACTTGATGGACATTGGTTTTTGTGTTTCATGATCATAATTTTCA  
 31281 AGCTCTGATTTTTTTCAGTTTCATCCATTGAGTAGGTATACCATCATGTCTTTTTTTTTTTTGTCTTTTTTTTTTTTTTTTTTTTGGAGCAGAG  
 31373 TCTTGCTCTGTGCGCCAGGCTGGAGTGCAGTGGTGAATCTCGGCTCACTGCAAGCTCCGCCTCCTGGGTTACACCATTCCTCCTGCCTCAG  
 31465 CCTCAGCCTCCCGAGTAGCTGGGACCACAGGTGCCACTACCACACCTGGCTAATTTTTTTGTATTTTTTTGTAGAGACGGGGTCTCACTGGG  
 31557 TTAGCCAGGATGGTCTCGATCTCCTGACCTGGTGAGCCGCCAGCCTCGGCCTCCCAAAGTGCTGGAATTACAGGCGTGAGCCACCGTGCCCG  
 31649 GCCCATGTCTTTGACCATTTGTTATAAACTATGTGTGTAACCTACTATAAACCATAGAAACCGATTATATAATAGCAACACTATTGTGAGTAA  
 31741 TAAGTGTATATAGCTTTTCCATATTTTATTCCGTTTCTTTGGATGCATTTATGATGTTTTTTTAAATAAGAGCATAACTTATTTATATGT  
 31833 TTACATTTTCTTTTAAAGAAGCTTGACTTCTCATCAGCTGAGCCAGAAGTGAAGTCATTTGAAGAGAAGTTTGGA AAAAGGATCCTTGTCAA  
 31925 GTGCAATGATTTATCTTTCAATTTGCAATGCTGTGTTGCCGAAAATGAAGAAGGACCCACTACAAATGTAATTTTTTCATTTTAAAAATAAAC  
 32017 ATTA AAAAAAAAAATAGGCAGAGGTTTCAGATGTACCTTTACAGTGCAGCCTGGATAAGAAATCCTAGTCCCTGGTATCAAAGAGGTGCAGTG  
 32109 TTTGGATCAGGATATGGAGGTTGTTAGCCTGCAAGGACAGGATGTTGCTGATGGAAGATGAGGGTGGCAGGTTTGTGCTCAGCTTTCAGGA

FIG. 12B  
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41401 CTTTTTCCCTTGCAATAATTACCCAGGGATATGTTCCAAGATTTAGTAAGAAAGCGATTCTGTCCGATAGATGATATTGCTAACATTTTATA  
 41493 AGAAGAGAGACTTGGTACTTTGTATTTGATTTGTTTCATGGTGGTATCTCATGGATAAGATGGTATCTCATCTTTTCCAACCTTCTGCAGGAAA  
 41585 TGCGAAGACATGAAGGCAAAGTATAAAAATAGAACGTTTTCTTTAAAACGTAGACCTTTTAAATGGTACTACGTTGGATAGTTTAGGTAATA  
 41677 ATACTACTAAAGTTTTTTCGTATGCAGCTTAATGTGTCTGTGTTTATTTGTACACTCATCTTCTTTGCATCCAGGTTTTACAGTCTTACCCC  
 41769 GATTTCGCTCTGGTTACACTGCACTCAAGCCAAGTAGGGCTGCTTGACTTTCTCTAAACCCACTGGGGACTTCCCTCTGCCATGCTTTTCTC  
 41861 TCTGCCCCAATTGTGTCCCCCTTCCTGCCTCATCAAGCAGCACATAAATCACAAACACATGCAGCATACACACTTCCCCTTTTCTTTGTCTT  
 41953 TCTCAGGGAACTCTACTCATCTTTCAAAGCCCAGTCTGTGGCTCACTTCTGTGCTGGGAGTCCTGGAGGCGGTACTTGGCTTCTCTGCCTG  
 42045 AGCCGCCTCCTCTTTTTAAGGGTGGATAATAACAGCCCCCTGCCCCCTAAACCGTGGTGGGGAATAAATGCAAAGGCATTAAGGTGATTTT  
 42137 TCCCACCATGAATACTGATCTCATCCCGTGTTCCTCTCGATAGATCTAGATACTCTGCCTTCTGGTAGAGGTTTGTACATACTCTGTGAAA  
 42229 GTGATTGCCCTCATATGCCGTAAAGTAGCTTACAGTGTCTACTGGACTTTTGGCTTCTTGAGGAAGAAATTATGTCTTGTGTTGCATTCTCTC  
 42321 ATGGTCCTGAGTACATACATTGCAGCATATCCTAAGCACTTGATAAATGCTTATTGAATTTTCTTCTTAGACATAAACTCAGTGGTTTTTGT  
 42413 TGAAACAAAAATATCTCAATTTCTTTCAATCATATATAGTTGTTTTTTTTTAAGTGACACCAAAGCTTTTAGGGAATATTTCTTTTCAAA  
 42505 AACACAGTTAGAAGATTAACTCACCACCAATAGCAGTCCAAACATACCTGTATTGCCAGCTAATCATTTTAAACGAGCCAATACAGGAAGTC  
 42597 AGGAAGGGAAGACCGGCTGCAGAAACACTTAGATAAGGACCCCAAATCTGTTGGCATGGGAGGACTGCTAGTTGATGATACCATTTCCCATTT  
 42689 CCTCTGTGGGAATTGTTGAGTCAGCAGAAATGGATGGGCAAGGAAATTTTCTTAAGAGAGAGTTTGAGCCTCACTTCTACATTC  
 42781 ACACAGAGACAGGAGCAGTTCCCAGAGGCCAGGCATCCTGCAAGTGTCTGTATTGCATGCTTACTTAATTCGTGTAATTTTAAGATGAGTTT  
 42873 TCATGTTCAAGGATTATTTTATAAATTTTGCATAGAATATAGGTACTCTTTAGCAAAACAAAGCAAAAAACCAAACTATTCTCAGTCATG  
 42965 AAAGAATTGAGTTTGTGTAACACGCACACAACCACCACTTTGGAAGTGCATAAAAAGGCAGTAAATCTTTATTGCCTGTGAGTGTTTGATG  
 43057 TCTAATAAACCAGATTCAACATAAACCATAAACTTTTGAATGGGTTTGAGATTGGGTTTTTAAAACTTAAAGCTGGCAAAAAAAAACA  
 43149 ACTTTTAAAAGCCCATGTGCTACATAATATGGAATAAATCAGAAATGTGCTTGGAAACACATGGAAAGAACGTCTTTACAGAAGCAGCAA  
 43241 CTAGAAGTAAATCTCTCAGCAGAGGGAGGAAATAGAATAAGAAATAACTATAGTTAGGCACAGAAGGACACAATACTATAGGAAGATTT  
 43333 CCAGTGAAGATCATTTAATTAAATATGTTGCTTAGAAACGTATTTTAATTGTGTTCCACCTCTCTCAAAAATTTATATGTGGAGGATGTTG  
 43425 GAGTGATCTTAAAAATGGTGATGAAGATGCCTGTTTCATTCATAGGTGGAATAATTAGGAGGGGGTGAATCCATTACCCTTGCATACTTAC  
 43517 TTATATTTAAAAGTATAATTTGTAATAAA

hCLASP4	-----MFPMEDISISVIGRQRRTVQ-----	20
hCLASP5	-----MTHLNSLDVQLAQELG-----	16
hCLASP3	-----MAERRAFAQKISRTVAAAEVRKQISGOYSGSPQLLKNLNIVG	41
hCLASP2	-----MLLFPYDDFQTAILRRQGRYICS-----	23
hCLASP7	-----MAASERRAFAHKINRTVAAAEVRKQVSRERSGSPHSSRRCSSSL	43
hCLASP1	MSFRGKVFVKREPSEFWKKRRTVRRVIQEEFHRFSSQEKPRLLEPLDYETVIEELEKTYRN	60
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hCLASP4	-----STVPEDA EKRAQSLFVKECIKTYSTDWHVVNYK	53
hCLASP5	-----DFT	19
hCLASP3	N-----ISHHTTVPLTEAVDPVDLEDYLITHPLAVDSGPLRDLIEFP	83
hCLASP2	-----TVPAKAE EEAQSLFVTECIKTYNSDWHLVNYK	55
hCLASP7	G-----VPLTEVVEPLDFEDVLLSRPPDAEPGPLRDLVEFP	79
hCLASP1	DPLQDLLFFPSDDFSAATVSWDIRTLYSTVPEDA EHKAENLLVKEACKFYSSQWHVVNYK	120
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hCLASP4	YEDFSGDFRMLPCKSLRPEKIPNHVFEIDEDCEKDED-----SSSLCSQKGGVIKQG	105
hCLASP5	DDDL DVVFTPKCRTLP-SLPEEGVELDPHVR-----DCVQTYIREWLI	63
hCLASP3	PDDIEVVYSPRDCRTLVS-AVPEE-SEMDPHVR-----DCIRSYTEDWAI	126
hCLASP2	YEDYSGEFRQLPNKVVKLDKLPVHVVEVDEEVDKDED-----AASLGSQKGGITKHG	107
hCLASP7	ADDLELLLQPRECRTEP-GIPKD-EKLDAQVR-----AAVEMYIEDWVI	122
hCLASP1	YEQYSGDIRQLPRAEYKPEKLPSHSFEIDHEDADKDEDTTSHSSSKGGGGAGGTGVFKSG	180
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hCLASP4	WLHKANVNSTIT--VTMKVFKRRYFYLTQLPDGSYILNSYKDEKNSKESK-GCIYLDACI	162
hCLASP5	VNRKNQGSPEIC--GFKKTGSRKDFHKT-LPKQTFESETLECSEPAAQA--GPRHLNVLC	118
hCLASP3	VIRKYHKLGTGF--NPNTLDKQKERQKG-LPKQVFESDEAPDGNSYQDDQDDLKRRSMSI	183
hCLASP2	WLYKGNMNSAIS--VTMRSFKRRFFHLIQLGDGSYNLNFYKDEKISKEPK-GSIFLDSCM	164
hCLASP7	VHRRYQYLSAAY--SPVTTDTQRRERQKG-LPRQVF EQDASGDERSGPEDSNDSRRGSGSP	179
hCLASP1	WLYKGNFNSTVNNTVTVRSFKKRYFQLTQLPDNSYIMNFYKDEKISKEPK-GCIFLDSC	239
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hCLASP4	DVVQCPKMRRHAFELKMLDKYSHYLAAETE QEME EWLITLKKIIQINTDSLVOEKKETVE	222
hCLASP5	DVSGKGPVTACDFDLRSLQPDKRLENLLQQVSAEDFEKQNEEARRTN-----RQAE	169
hCLASP3	DDTPRGSWACSI FDLKNSLPDALLPNLLDRTPNEEIDRQNDQORKSN-----RHKE	234
hCLASP2	GVVQNNKVRRAFELKMQDKSSYLLAADSEVEME EWITILNKILQLN-----FEAAMQEK	219
hCLASP7	EDTPRSSGASSI FDLRNLAADSLPSLLERAAPEDVDRRNETLRRQH-----RPPA	230
hCLASP1	GVVQNNRLRKYAFELKMNDLTYFVLAAETESDMDEWIHTLNRILQISPEGPLQGRSTEL	299
* : : . : : :		
hCLASP4	TAQDDETSS----QGAENIMASLERSMHP ELMKYGRETEQLNKL SRGDGRQNLFSFDSE	278
hCLASP5	LFALYPSVD----EEDAVEIRPVPECPKEHLG-----N-----RILVKLLTLKFEIE	212
hCLASP3	LFALHPSPD----EEEPIERLSVPDIPKEHFG-----QRLLVKCLSLKFEIE	277
hCLASP2	RNGDSHEDD----EQSKLEGSGSGLDSYLP ELAKSAREAEIK---LKSESRVKLFYLDPD	272
hCLASP7	LLTLYPAPD----EDEAVERCSRPEPPREHFG-----QRILVKCLSLKFEIE	273
hCLASP1	TDLGLDSL DNSVTCECTPEETDSS ENNLHADFAKYLTETEDTVKTRNMERLNLFSLDPD	359
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hCLASP4	VQRLDFS----GIEPDIKP-FEEKCNKRFLVNCHDLTFNILGQIGDNAKG PPTNVEPFFI	333
hCLASP5	IEPLFAS----IALYDVKERKKISENFHCDLNSDQFKGFLRAHTPSVAASSQARS AVFSV	268
hCLASP3	IEPIFAS----LALYDVKEKKKISENFYFDLNS EQMKGLLRPHVPPAAITTLARSAIFSI	333
hCLASP2	AQKLDFS----SAEPEVKS-FEEKFGKRILVKCNDSL FNLQCCVAENE EGPTTNVEPFFV	327
hCLASP7	IEPIFGI----LALYDVREKKKISENFYFDLNSDSMKGLLR AHGTHPAISTLARSAIFSV	329
hCLASP1	IDTLKLQKKDLLEPESVIKPFEEKAAKRIMIICKALNSNLQGCVTENENDPITNIEPFFV	419
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FIG. 13  
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hCLASP4	NLALFDVKNNCKISADFHVDLNPPSVREMLWGSSTQLASDGSP---	KGSSPESYIHGIAE	390
hCLASP5	TYPSSDIYLVVKIEKVLQOQD----	IGDCAEPYTVIKESDG-----	GKSKE-KIEKLKL 317
hCLASP3	TYPQDVFLVIKLEKVLQOQD----	IGECAEPYMI FKEADA-----	TKNKE-KLEKLKS 382
hCLASP2	TLSLFDIKYNRKISADFHVDLNHFSVRQMLATTSPALMNGS-----	GQSPSVLKGI LHE	381
hCLASP7	TYPSPDIFLVIKLEKVLQOQD----	ISECCEPYMVLKEVDT-----	AKNKE-KLEKLRL 378
hCLASP1	SVALYDLRDSRKISADFHVDLNHAAVRQMLLGASVALENGNIDTITPRQSEEPHIKGLPE		479
	. . * : * : . : : . : : . . . :		
hCLASP4	SQLRYIQQGI FSVTNPHPEIFLVARIEKVLQGNITHCAEPYIKNSDPVKTAQKVHRTAKQ		450
hCLASP5	QAESFCQR-----	LGKYRMPFAWAPISLSSFFNVSTLEREVDVDSVVGRSPVGERRTLA	372
hCLASP3	QADQFCQR-----	LGKYRMPFAWTAIHLMNIVSSAGSLERDSTEVEISTGERKGSWSERR	437
hCLASP2	AAMQYPKQGI FSVTCPHPDI FVARIEKVLQGSITHCAEPYMKSSDSSKVAQKVLKNAKQ		441
hCLASP7	AAEQFCTR-----	LGRYRMPFAWTAVHLANIVSSAGQLDRDSD----	SEGERRPAWTD RR 429
hCLASP1	EWLKF PKQAVFSVSNPHSEIVLVAKIEKVLGMN IASGAEPYIKNPDSNKYAQKILKSNRQ		539
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hCLASP4	VCSRLGQYRMPFAWAARPI FKDTQGS LDDLGRFSPLYKQDSSKLSS EDILKLLSEYKKPE		510
hCLASP5	QSRRLSERALSLEENG VGSNFKTS-----	TL SVSSFFKQEGDRLSDEDLFKFLADYKRSS	427
hCLASP3	NSSIVGRRSLERTTSGDDACNLTSFR-PATLITV TNFFKQEGDRLSDEDLYKFLADMRRPS		496
hCLASP2	ACQRLGQYRMPFAWAARTLFKDASGNLDKNARFSAIYRQDSNKLSNDDMLKLLADFRKPE		501
hCLASP7	---RRGPQ--	DRASSGDDACSFSGFR-PATLITV TNFFKQEAERLSDEDLFKFLADMRRPS	483
hCLASP1	FCSKLGKYRRAFAWAVRSVFKDNQGNVDRDSRFSPLFRQESSKISTEDLVKLVSDYRRAD		599
	. : : : * : : : * : * : : : : . . .		
hCLASP4	--KTKLQIIPGQLNITVECV PVDLSNCITSSYVPLKPFE-KNCQNITVEVEEFVPEMTKY		567
hCLASP5	SLQRRVKSIPGLLRLEISTAPEI INCCLTPEMLPVKFPF-ENRTRPHKEILEFP--	TREV	484
hCLASP3	SVLRRLRPITAQLKIDISPAPENPHYCLTPELLQVKLYP-DSRVRPTREILEFP--	ARDV	553
hCLASP2	K-MAKLPVILGNLDITIDNVSSDFPNYVNSSYIPTKQFETCSKTPITFEVEEFVPCIPKH		560
hCLASP7	SLLRRLRPVTAQLKIDISPAPENPHFCLSPELLHIKPYP-DPRGRPTKEILEFP--	AREV	540
hCLASP1	R-ISKMQTIPGSLDIAVDNVPLEHPNCVTSSFI PVKPFNMMAQTEPTVEVEEFVYDSTKY		658
	: : : . * : : . . : : : : * : : * : * : .		
hCLASP4	CYPFTIYKNHLYVYPLQLKYDSQKTFAKARNIAVCVEFRDSDES DASALKCIYGKPAAGSV		627
hCLASP5	YVPHTVYRNLLYVYPQRLNFVN--	KLASARNITIKIQFMCG-EDASNAMPVIFGKSSGPE	541
hCLASP3	YVPNTTYRNLLYIYPQSLNFAN--	RQGSARNITVKVQFMYG-EDPSNAMPVIFGKSSCSE	610
hCLASP2	TOPYTIYTNHLYVYPKYLYDSQKSF AKARNIAICIEFKDSDEEDSQPLKCIYGRPGGPV		620
hCLASP7	YAPHTSYRNLLYVYPHSLNFSS--	RQGSVRNLAVRVQYMTG-EDPSQALPVIFGKSSCSE	597
hCLASP1	CRPYRVYKNQIYIYPKHLKYDSQKCFNKARNITVCIEFKNSDEESAKPLKCIYGKPEGPL		718
	* * * : : * : : . . * : : : : : : . * . : : : * : : .		
hCLASP4	FTTNAYAVVSHHNQNP EFYDEIKIELPIHLHQKHLLFTFYHVSC	INTKGTTKKQDTVE	687
hCLASP5	FLQEVYTAVTYHNKSPDFYEEVKIKLP AKLTVNHHLLFTFYHISCOQ-----	KQGASVE	595
hCLASP3	FSKRAYTAVVYHNRS PDFHEEIKVKLPATLTDHHL LFTFYHVSCQ-----	KQNTBLE	664
hCLASP2	FTRSAFAAVLHHHQNP EFYDEIKIELPTQLHEKHLLLTFFHVSCDNSSKGSTKKRDVVE		680
hCLASP7	FTRSAFTPVVYHNKSPEFYEEFKLHLPACVTENHLLFTFYHVSCQ-----	RPGTALE	651
hCLASP1	FTSAAYTAVLHHSQNPDFSDEVKIELPTQLHEKHHLIFS FYHVTCDINAKANAKKKEALE		778
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hCLASP4	TPVGFAWVPLLKDGRITFEQQLPV SANLPPGYLNLNDAESRRQCNVDIKWVDGAKPLLK		747
hCLASP5	TLLGYSWLPILLNERLQTGSYCLPVALEKLPPNYSMHSAEKVPLQNPPIKWAEGHKGVFN		655
hCLASP3	TPVGYTWIPMLQNGRLKTGQFCLPVSL EKPPQAYSVLSPEVP---	LPGMKWVDNHKG VFN	721
hCLASP2	TQVGYSWLPPLLKDGRVVTSEQH IPV SANLPSGHLGYQELGMGRHYGPEIKWVDGGKPLLK		740
hCLASP7	TPVGFTWIPLLQHGRRLRTGPFCLPVSV DQPPPSYSVLTPDVA---	LPGMRWVDGHKG VFS	708
hCLASP1	TSVGYAWLP LMKHDQIASQEYNIP IATSLPPNYLSFQDSASGKHGGS DIKWVDGGKPLFK		838
	* : : : * : : : . : : : : * : : : : : : : : : : : : : : : *		

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hCLASP4	FKSHLESTIYTQDLHVHKFFHHCQLIQS-----GSKEVPGELIKYLKCLHAM	794
hCLASP5	IEVQAVSSVHTQDNHLEKFFTLCHSLESQVTFPIRVLDQKISEMALEHELKLSIICLNSS	715
hCLASP3	VEVVAVSSIHTQDPYLDKFFALVNALDEH-LFPVRIGDMRIMENNLENELKSSISALNSS	780
hCLASP2	ISTHLVSTVYTQDQHLHNFFQYCKTES-----GAQALGNELVKYLKSLHAM	787
hCLASP7	VELTAVSSVHPQDPYLDKFFTLVHVLEEG-AFPFRLKDTVLESEGNVEQELRASLAALRLA	767
hCLASP1	VSTFVVSTVNTQDPHVNAFFQECQKREK-----MSQSPTS NFIRSCKNLLNVE	887
	.. *:: ** ::. ** : ..	
hCLASP4	EIQVMIQFLPVILMQLFR-----VLTNMTH-----EDDVP	824
hCLASP5	RLEPLVLFLHLVLDKLFQLSVQPMVIAGQTANFSQFAFESVVAIANSLHNSKDLSDQH	775
hCLASP3	QLEPVVRFLHLLLDKLI LLVIRPPVIAGQIVNLGQASFEAMASIINRLHKNLEGNHDQH	840
hCLASP2	EGHVMIAFLPTILNQLFR-----VLT-RAT-----QEEVA	816
hCLASP7	SPEPLVAFSHHVLDKLVRLVIRPPIISGQIVNLGRGA FEAMAHVVS LVHRSLEAAQDARG	827
hCLASP1	KIHAIMSFLPIILNQLFK-----VLVQNE-----EDEIT	916
	. : * : * : *	
hCLASP4	INCTMV-LLHIVSKCHEEGLDS-----YLRSEFIKYS-----FRPEKP	860
hCLASP5	RNCLLASVHYVFRLEPEVQRDVPKSGAPTALLDPRSHTYGR TSAAAVSSKLLQARVMSS	835
hCLASP3	RNSLLASYIHVFRLPNTYPNSSSPG-PGGLGGSVHYATMARS AVR PASLNLNRSRSLN	899
hCLASP2	VNVTRV-I IHVVAQCHEEGLES-----HLRSYVKYA-----YKAEPY	852
hCLASP7	HCPQLAAYVHYAFRLPGTEPSLPDGAPP----VTVQAATLARGSGRPASLYLARSKSISS	883
hCLASP1	TTVTRV-LPDI VAKCHEEQLDH-----SVQSYIKFV-----FKTRAC	952
	. . . :	
hCLASP4	SAPQAQLIH-----ETLATTMIAILKQS-----	883
hCLASP5	SNPDLAGTHSAADEEVKNIMSSKIADRNC SRMSYYCSGSSDAPSSPA-----	882
hCLASP3	SNPDISGTPTSPDDEVRSIIGSKGLDRSNSWVNTGGPKAAPWGSNPSPSAESTQAMDRSC	959
hCLASP2	VASEYKTVH-----EELTKSMTTILKPS-----	875
hCLASP7	SNPDLAVAPGSVDDEVSRILASKLLHEELA-LQ-----	915
hCLASP1	KE---RPVH-----EDLAKNVTGLLKS-----	972
	: . . .	
hCLASP4	-----ADFLSINKLLKYS-----WFFFEIIAKSM	907
hCLASP5	-----APRPASKKHFEELALQ-----MVVSTGMVKSM	910
hCLASP3	NRMSHTETSSFLQTLTGRLPTKKLFHEELALQWVVC SGSVRESALQQA WFFFELMVKSM	1019
hCLASP2	-----ADFLTSNKLLRYS-----WFFFDVLIKSM	899
hCLASP7	-----WVVS S AVREAILQHA-----WFFFQLMVKSM	942
hCLASP1	-----DSPTVKHVLKHS-----WFFFAILKSM	995
	. * .. : ***	
Cadherin Cleavage		
hCLASP4	ATYLLEENKIKL RRGQRFPE TYHHVLHSLLLAIIPHVTIRYAEIPDE---SRNVNYSLAS	964
hCLASP5	AQHVNMDKRDSE RRTFRSDFMDDITTIVNVVTSEIAALLVKPQKENEQA EKMNISLAF	970
hCLASP3	VHHLYFNDKLEA RRSRFRPFMDDIAALVSTIASDIVSRFQKDTM---VERLNTSLAF	1076
hCLASP2	AQH LIENSKVKLLRNQRF PASYHHAETVVNMLMPHITQKFGDNPEA---SKNANHSLAV	956
hCLASP7	ALHLLLGQRLDTERKLRFPGRFLDDIT ALVGSVGLVITRVHKDVEL---AEHLNASLAF	999
hCLASP1	AQH LI DTNKIQLRPQRFPE SYQNELDNLMVLS DHVIWKYKDALEE---TRRATHSVAR	1052
	. : . : . * *. : . : : . . . . . *	
hCLASP4	FLKRCLTLMDRGFI FNLINDYISGFSPKDP-----KVLAEYKFEFLQTICNHEHYIPLNL	1019
hCLASP5	FLYDLLSLMDRGFVENLIRHYCSQLSAKLSNL---ETLISMRLFLRILCSHEHYLNLNL	1027
hCLASP3	FLNDLLSVMDRGFVFS LIKSCYKQVSSKLYSLPNPSVLVSLRLD FLRIICSHEHYVTNLNL	1136
hCLASP2	FIKRCFTFMDRGFVEKQINNYISCFAPGDP-----KTLFEYKFEFLRVVCNHEHYIPLNL	1011
hCLASP7	FLSDLLSLVDRGFVFS LVRAHYKQVATRLQSSPNPAALLTLRMEFTRILCSHEHYVTNLNL	1059
hCLASP1	FLKRCFTFMDRGCVFKMVNNYISMFS S GDL-----KTL CQYKFDFLQEV CQHEHFIPCL	1107
	*: :::*** :*. . . . . *	

FIG. 13  
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Cadherin EC motif

hCLASP4 PMAFAKPKLQR-----VQDS--NLEYSLSDEYCKHHFLVGI LLRETSI 1060

hCLASP5 FFMNADTAPTSP--CPSISSQNSSSCSSFQDQKIASMFDLTSEYRQQHFLTGI LFTELAA 1085

hCLASP3 PCSLLTPPASPSPSVSSATSQSSGFSTNVQDQKIANMFELSVPFRRQQHYLAGIVLTELAV 1196

hCLASP2 PMPFGKGRIQR-----YQDL--QLDYSLTDEFECRNHFLVGI LLREVGT 1052

hCLASP7 PCCPLSPPASPSPSVSSTTSQSSTFSSQAPDPKVTSMFELSGPFRQQHFLAGI LLTELAL 1119

hCLASP1 PIRSANIPDPLTP-----SES----TQELHASDMPEYSVTNEFCRKHFLIGI LLREVGF 1157

. : : : : \* \* \* \* \*

hCLASP4 ALQDN----YEIRYTAISVIKNLLIKHAFDTRYQHKNQQAKIAQLYLPFVGLLLENIORL 1116

hCLASP5 ALDAEGEGISKVQRKAVSAIHSLLSSHDLDPKVCPEVKVKIAALYLPVGI ILDALP-- 1143

hCLASP3 ILDPDAEGLFGLHKKVINMVHNLSSHDSDPYSDPQIKARVAMLYLPLIGI IMETVP-- 1254

hCLASP2 ALQEFR----EVRLIAISVLKNLLIKHSFDDRYASRSHQARIATLYLPLFGLLIENVQRI 1108

hCLASP7 ALEPEAEGAFILHKKAISAVHSLLCGHDTDPYAEATVKARVAELYLPLLSIARDTLP-- 1177

hCLASP1 ALQEDQ----DVRHLALAVLKNLMAKHSFDDRYREPRKQAQIASLYMPLYGMLLDNMPRI 1213

\* : : : : \* \* \* : : \* \* \* \* : : :

hCLASP4 AGRDTLYSCA-----AMPN-S----ASRDEFPCGFTSPANRGSLSLTDKDTAYGS 1160

hCLASP5 -----QL-----CDFTVADTRRYRTSGSD----- 1162

hCLASP3 -----QLY-----DFTETHNQGRPICIAATDD-- 1276

hCLASP2 NVRDVSPFPVNAGMTVKDESLALPA-VNPLVTPQKGSTLDNSLHKDLLGAISGIASPYTT 1167

hCLASP7 -----RLH-----DFAEGPGQRSRLASMLDSLTE 1201

hCLASP1 YLKDLYPFTVNTSNQGSRDDLSLNGGFQSQTAKHANSVDTSFSDVLNSIAAFSSIAIS 1273

. :

hCLASP4 FQ-NGHGIKREDSRGS LIPEGATGFPDQNGTGEN-----TRQSSTRSSVSQYNRLDQYE 1213

hCLASP5 -----EEQEGAGAINQNVALAIAGNNFNLKT-----SGIVLSSLPYKQYNMLNADT 1208

hCLASP3 -----YESESGSMISQTVAMAIAGTSVPQLTR----PGSFLTSTSGRQHTTFSAES 1324

hCLASP2 STPNINSVRNADSRGSLISTDSGNSLPERNSEKSNSLDKHQSSSTLGNSVVRCDKLDQSE 1227

hCLASP7 -----GEGDIAGTINPSVAMAIAGGPLAPGSR----ASISQGPPTASRAGCALSAES 1249

hCLASP1 -----TVNHADSRASLASLDSNPSTNEKSSEKTDNCEKIPRPLALIGSTLRFDRLDQAE 1327

. : : : . . : :

hCLASP4 IRSLLMCYLYIVKMISED TLLTYWNKVSPQELINILILLEVCLFHFYRMGKRNIARVHDA 1273

hCLASP5 TRNLMICFLWIMKNADQSLIRKWIADLPSTQLNRILDLLFICVLCFEYKKGKQSSDKVSTQ 1268

hCLASP3 SRSLICLLWVLKNADET V LQKWFDTLSVLQLNRLLDLLYLCVSCFEYKKGKVFERMNSL 1384

hCLASP2 IKSLLMCFLYILKSMSDDALFTYWNKASTSELMDFFTISEVCLHQFQYMGKRYIARNQEG 1287

hCLASP7 SRTLLACVLWVLKNT EPALLQRWATDLTLPQLGRLLDLLYLCLAAFEYKKGKKA FERINSL 1309

hCLASP1 TRSLLMCFLHIMKTISYETLIAYWQRAPSPESVDFFSILDVCLQNFYLGKRNIIRKIAA 1387

: . \* \* : \* . : : . : : : \* : \* \* \* : :

hCLASP4 WLSKHFGIDR-----KSQTMPALRNRSQVMQARLQHLSSLESS----- 1311

hCLASP5 VLQKSRDVKAR-----LEEALLRGE GARGEMRRRAPGNDRFPGLNEN--- 1311

hCLASP3 TFKKSKDMRAK-----LEEAILGSIGARQEMVRRSRGQLERSPSGSAFGSQ 1430

hCLASP2 LGPIVHDRKS-----QTLPVSRNRTGMMHARLQQLGSLDNS----- 1323

hCLASP7 TFKKSLDMKAR-----LEEAILGTIGARQEMVRRSRERSPFGNPEN----- 1350

hCLASP1 AFKEVQSTQNNGT LKGSNPSCQTSGLLAQWMHSTSRHEGHKQHRSTLPIIRGKN----- 1442

. : :

hCLASP4 -----FTLNHSSTTTEADIFHQALLEGNTATEVSLTVLDTISFFTQCFTQLL 1359

hCLASP5 --LRWKKEQTHWRQANEKLDKTKAE L DQEALISGNLATEAHLIILDMQENIIQASS-ALD 1368

hCLASP3 ENLRWRKDMTHWRQNT EKLDKSRAEIEHEALIDGNLATEANLIILDTLEIVVQTVS-VTE 1489

hCLASP2 -----LTFNHSYGHSDADV LHQS LLEANIATEVCLTALDTLSLFTLAFKNQLL 1371

hCLASP7 --VRWRKSVTHWKQTS DRVDKTKDEMEHEALVEGNLATEASLVVLDLTLEIIVQTVM-LSE 1407

hCLASP1 --ALSNPKLLQMLDNTMTSNEI DIVHHVDTEANIATEGCLTILDVSLFTQTHQRQLQ 1500

. . : : . \* \* \* \* \* . .

FIG. 13  
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hCLASP4 NNDGHNPLMKKVFDIHLAFLKNGQSEVSLKHVFASLRAFISKFPSAFFKGRVNMCAAFCY 1419  
hCLASP5 CKDS---LLGGVLRVLVNSLNCDQSTTYLTHCFATLRALIAKFGDLLFEEVEQCDFDLCH 1425  
hCLASP3 SKES---ILGGVLKVLLHSMACNQSAVYLQHC FATQRALVSKFPPELLFEEETEQCADLCL 1546  
hCLASP2 ADHGHNPLMKKVFDVYLCFLQKHQSETALKNVFTALRS LIYKFPSTFYEGRADMCAALCY 1431  
hCLASP7 ARES---VLGAVLKVVLYSLGSAQSALFLQHGLATQRALVSKFPPELLFEEDTELCADLCL 1464  
hCLASP1 QCDCQNSLMKRGFDTYMLFFQVNO SATALKHVFASLRFLVCKFPSAFFQGPADLCGSFCY 1560  
. : : : : \* \* \* : : : \* : : \* \* . : : : . : \* : \*

hCLASP4 EVLKCCTSKISSTRNEASALLYLLMRNNFEYTKRKTFLRTHLQII IAVSqliADVALSGG 1479  
hCLASP5 QVLHHCSSSMDVTRSQCACATLYLLMR--FSFGATSNFARVKMQVTMSLASLVGRAPDFNE 1483  
hCLASP3 RLLRHCS SIGTIRSHPSASLYLLMR--QNFEIGNNFARVKMQVPMSLSSLVGTSQNFNE 1604  
hCLASP2 EILKCCNSKLSSIRTEASQLLYFLMRNNFDYTGKKS FVRTHLQVIISVSLIADVVGIGE 1491  
hCLASP7 RLLRHCGSRISTIRTHASASLYLLMR--QNFEIGHNFARVKMQVTMSLSSLVGTTQNFSE 1522  
hCLASP1 EVLKCCNHRSRSTQTEASALLYLFMRKNFEFNKQKSIVRSHLQLIKAVSqliADAG-IGG 1619  
.: : \* : . . . . \* : : \* : . : \* : : : : : : .

hCLASP4 SRFQESLFIINNFANS DRPMKATAFP AEVKDLTKRIRTVLMATAQMKEHEKDPEMLIDLQ 1539  
hCLASP5 EHLRRSLRTILAYSEEDTAMQMTFPTQVEELLCNLNSILYDTVKMREFQEDPEMLMDLM 1543  
hCLASP3 EFLRRSLKTILTYAEEDLELRETTFPDQVQDLVFNLMILSDTVKMKEHQEDPEMLIDL 1664  
hCLASP2 TRFQQSLSIINNCANS DRLIKHTSFSSDVKDLTKRIRTVLMATAQMKEHENDPEMLVDLQ 1551  
hCLASP7 EHLRRSLKTILTYAEEDMGLRDSTFAEQVQDLMFNLHMILTDTVKMKEHQEDPEMLIDL 1582  
hCLASP1 SRFQHS LAITNNFANGDKQMKNSNFPAEVKDLTKRIRTVLMATAQMKEHEKDPEMLVDLQ 1679  
.: : \* : : \* : : : \* : : : \* : : : \* : : : \* : : : \* : : : \*

transmembrane  
hCLASP4 YSLAKSYASTPELRKTWLD SMAKIHVKNGD FSEAAMCYVHVAALVAEFLHRKK----- 1592  
hCLASP5 YRIAKSYQASPDRLRLTWLQNM AEKHTKKKYTEAAMCLVHAAALVAEYLSMLEDH----- 1598  
hCLASP3 YRIAKGYQTSPE-RLTWLQNMAGKHSERSN HAEAAQCLVHSAALVAEYLSMLEDR----- 1718  
hCLASP2 YSLAKSYASTPELRKTWLD SMARIHVKNGLSEAAMCYVHVTALVAEYLTRKG----- 1604  
hCLASP7 YRIARGYQGPSDLRLTWLQNMAGKHAELGN HAEAAQCMVHAAALVAEYI ALLEDQ----- 1637  
hCLASP1 YSLANSYASTPELRRTWLESMAKIHARNGDLSEAAMCYIHIAALIAEYI KRKGYWKVEKI 1739  
\* : \* . \* : : \* \* \* : . \* \* : \* \* : \* : \* : \*

-----LFPNGCSAFKKITPNIDE EGAMKEDAGMMD----- 1622  
-----SYLPVGSVSFQNISSNVLEESV VSED TLSPDEGVD 1633  
-----KYLPGCVTFQNISSNVLEESA VSDDVSPDEEGI 1753  
-----VFRQGCTAFRVITPNIDE EASMMEDVGMQD----- 1634  
-----RHLPVGCVSFQNISSNVLEESA ISDDILSPDEEGF 1672  
CTASLLSE DTHPCDSNSLLTTPSGGSMFSGMGP AFLSITPNIKEEGA AKEDSGMHD----- 1795  
: \* : \* : : \* : : \* : \*

ITAM  
hCLASP4 ---VHYSEEVLLELLEQCVDGLWKAERYEII SEISKLI VPIYEKRREFEKLTVYRTLHG 1679  
hCLASP5 CAGQYFTESGLVGLLEQA AELFSTGGLYETVNEVYKLVIPILEAHREFRKLTLTHSKLQR 1693  
hCLASP3 CSGKYFTESGLVGLLEQA AASFMSMAGMYEAVNEVYKVLIP IHEANRDAKKLSTIHGKLQE 1813  
hCLASP2 ---VHFNE DVLME LLEQCADGLWKAERYELIADIYKLIPIIYEKRR----- 1677  
hCLASP7 CSGKHFTELGLVGLLEQA AGYFTMGGLYEAVNEVYKNLIP ILEAHRDYKKLA AVHKGKLQE 1732  
hCLASP1 ---TPYNE NILVEQLYMCGEFLWK SERYELIADV NKPIIAVFEKQ RDKKLS DLYYDIHR 1852  
.: \* \* : \* . : . \* \* : : : \* : : : \* : \*

ITAM DOCK motif DOCK motif ITAM  
hCLASP4 AYTKILEVMHTKKRLLGTFFRVA FYGQSFFEEEDGKEYIYKEPKLTGLSEISLR LVKIY 1739  
hCLASP5 AFDSIVNKDH--KRMFGTYFRVGF FG-SKFGDLDEQEFVYKEHAITKLPEISHRLEAFY 1750  
hCLASP3 AFSKIVHQSTGWERMFGTYFRVGF FG-TKFGDLDEQEFVYKEHAITKLAEISHRLEGEFY 1872  
hCLASP2 -----DFFEDEDGKEYIYKEPKLTPLSEISQRLLKIYS 1710  
hCLASP7 AFTKIMHQSSGWERVFGTYFRVGF FG-AHFGDLDEQEFVYKEPSITKLAEISHRLEEEYT 1791  
hCLASP1 SYLKVAEVVNSEKRLFGRYRVA FYGQGFFEEEGKEYIYKEPKLTGLSEISQRLLKIYA 1912  
\* : : : : \* : : \* : \* : \*

FIG. 13  
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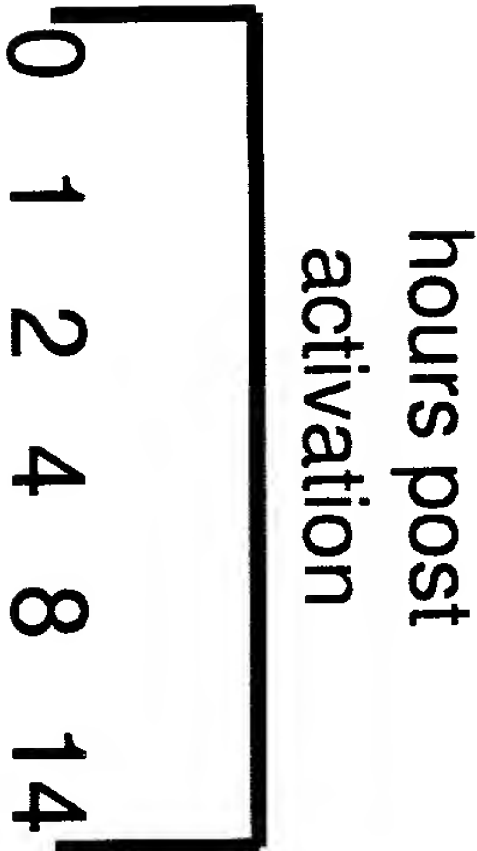
	ITAM	ITAM	
hCLASP4	EKFGTENVKIIQDSDKVNAKELDPHYAHIQVTHVKE	YFDDKELTERKTEFERNHNISR	FV 1799
hCLASP5	QCFGAEFVEVIKdstpvdktklDPNKAYIQITFVE	YFDEYEMKDRVTYFEKNFNLRR	FM 1810
hCLASP3	ERFGEDVVEVIKDSNPVDKCKLDPNKAYIQITFVE	YFDYEMKDRITYFDKNYNLRR	FM 1932
hCLASP2	DKFGSENVKMIQDSGKVNPKDLDSKYAYIQVTHV	IEFFDEKELQERKTEFERSHN	IRRM 1770
hCLASP7	ERFGDDVVEIIKDSYPVDKSKLDSQKAYIQITFVE	YFDYELKDRVTYFDRNYGLRT	FL 1851
hCLASP1	DKFGADNVKIIQDSNKNVNPKDLDPHYAYIQVTHV	TEFFEEKEIEDRKTDFEMHHN	INRFV 1972
	: ** : *::** * : .***: *::** : *:: * : : * * : ... *		
		ITAM	DOCK motif
hCLASP4	FEAPYTLGKKQGCIEEQCKRRTILTTSNSFPYVKKR	IPINCEQQINLKPIDGATDEIK	D 1859
hCLASP5	YTPFTLEGRPRGELHEQYRRNTVLTTHAFPIKTR	ISVIQKEEFVLTPIEVAIEDMK	K 1870
hCLASP3	YCTPFTLDGRAHGEHQFKRKTILTTSNHFPIKTR	VNVTHKEEILTPIEVAIEDMQ	K 1992
hCLASP2	FEMPFTQTGKRQGGVEEQCKRRTILTAIHCFPIV	KRIPVMYQHHTDLNPIEVAIDEM	SK 1830
hCLASP7	FCTPFTPDGRAHGEHQHKKRKTLLSTDHAFPIKTR	IRVCHREETVLTPEVAIEDMQ	K 1911
hCLASP1	FETPFTLSGKKHGGVAEQCKRRTILTTSNHFPIV	KRIQVISQSSTELNPIEVAIDEM	SR 2032
	: ** * : * : * : * : *::* : *::** : * : . *::* : :		
	Coiled-coil		
hCLASP4	KTAELQKLCSSSTDVDMIQLQLKLQGVWSVQVNAG	PLAYARAFLNDSQASKYPPKKVSEL	K 1919
hCLASP5	KTQLAVAINQEPDAKMLQMVLC	SVGATVNOGPLEVAQVFLAEIPADPKLYRHHNKL	R 1930
hCLASP3	KTQELAFATHQDPADPKMLQMVLC	SVGTTVNOGPLEVAQVFLSEIPSDPKLFRHHNKL	R 2052
hCLASP2	KVAELRQLCSSAEVDMIKLQLKLQGVWSVQVNAG	PLAYARAFLDDTNTKRYPDNKVKLLK	1890
hCLASP7	KTRELAFAEQDPPDAKMLQMVLC	SVGPVNOGPLEVAQVFLAEIPEDPKLFRHHNKL	R 1971
hCLASP1	KVSELNQLCTMEEVDMISLQLKLQGVSVSVKNAG	PMAYARAFLEETNAKKYPDNOVKLLK	2092
	*. : * * ** : *** * . ** ** : *::** : . : . *		
	Coiled-coil		
hCLASP4	DMFRKFIQACSALELNERLIKEDQVEYHEGLKSNFR	DMVKELSDIHEQILQEDTMHSP	1979
hCLASP5	LCFKEFIMRCGEAVEKNKRLITADQREYQOELKKNY	NKLKENLRPMIERKIPELYKPIFR	1990
hCLASP3	LCFKDFTKRCEDALRKNKSLIGPVQKEYQRELGLSSP	-----	2090
hCLASP2	EVFRQFVEACGOALAVNERLIKEDQLEYQEEMKANY	REMAKELSEIMHEQICPLEEKTS-	1949
hCLASP7	LCFKDFCKKCEDALRKNKALIGPDQKEYHRELERNY	CRLREALQPLLTQRLPQLMAPTP-	2030
hCLASP1	EIFRQFADACGOALDVNERLIKEDQLEYQEELRSHY	KDMLSELSTVMNEQITGRDDLSCR	2152
	*:. * * : * : * * * *::* :		
	PDZ ligand		
hCLASP4	WMSNTLHVFCASISGTSSDRGYGSPRYAEV	--	2008
hCLASP5	VESQKRDSFHRSSFRCETQLSQGS	-----	2015
hCLASP3	-----		
hCLASP2	VLPNSLHIFNAISGTPTSTMVHGMTSSSSVV		1980
hCLASP7	--PGLRNSLNRASFRKADL	-----	2047
hCLASP1	GVDQTCTRVISKATPALPTVSISSSAEV	--	2180

FIG. 13  
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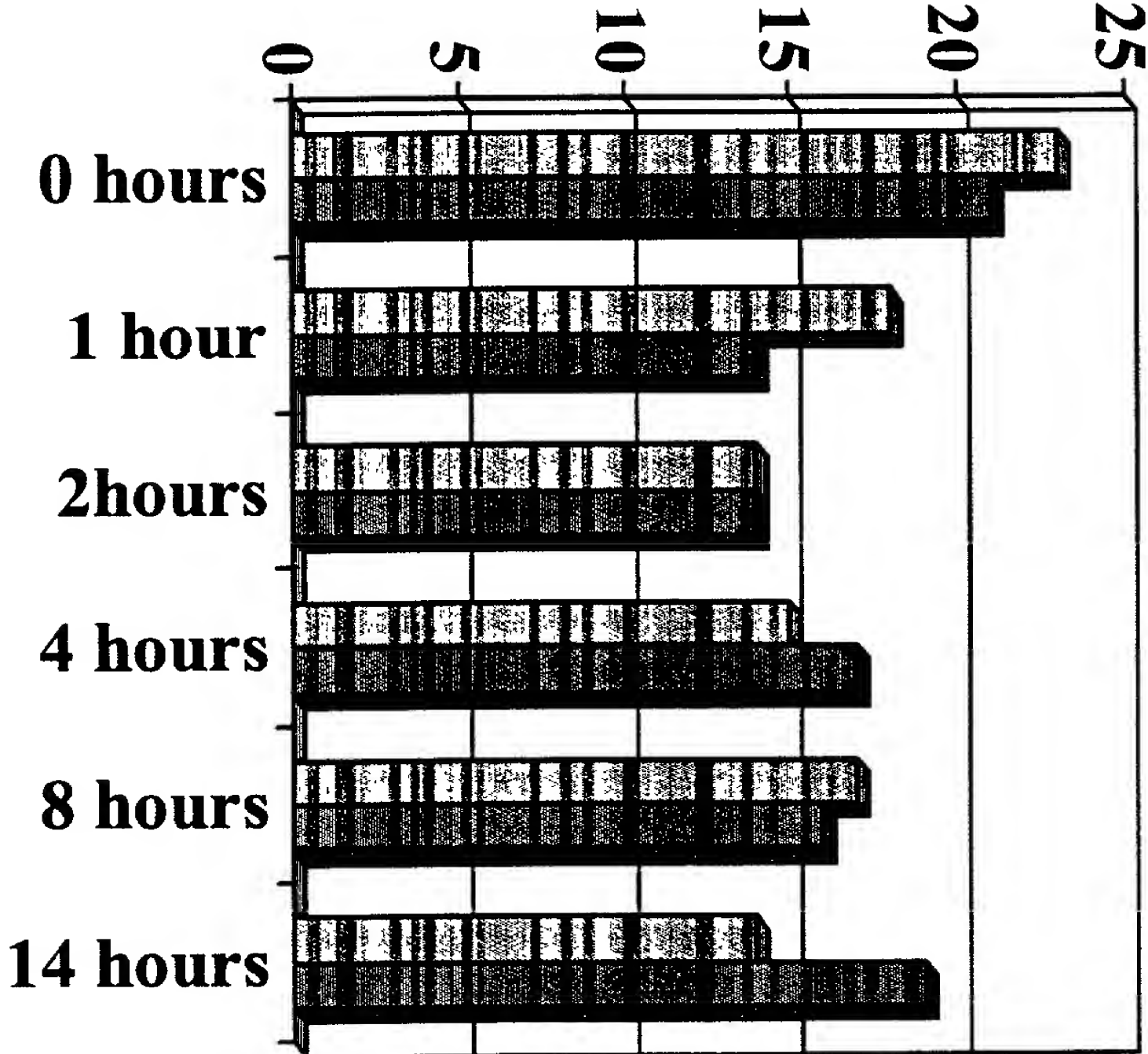
# Human CLASP-2 expression in T cells upon activation

**A)**



**B)**

% of total signal



CLASP-2 (A)  
CLASP-2 (B)

~ 7.5 kb -

CLASP-2

2.5 kb -



28s rRNA Ethbr. staining

09687837 101300

FIG. 14

**DECLARATION AND POWER OF ATTORNEY**

As a below named inventor, I declare that:

My residence, post office address and citizenship are as stated below next to my name; I believe I am the original, first and sole inventor (if only one name is listed below) or an original, first and joint inventor (if plural inventors are named below) of the subject matter which is claimed and for which a patent is sought on the invention entitled: **CLASP-2 TRANSMEMBRANE PROTEINS** the specification of which is attached hereto.

I have reviewed and understand the contents of the above identified specification, including the claims, as amended by any amendment referred to above. I acknowledge the duty to disclose information which is material to patentability as defined in Title 37, Code of Federal Regulations, Section 1.56. I claim foreign priority benefits under Title 35, United States Code, Section 119 of any foreign application(s) for patent or inventor's certificate listed below and have also identified below any foreign application for patent or inventor's certificate having a filing date before that of the application on which priority is claimed.

**Prior Foreign Application(s)**

Country	Application No.	Date of Filing	Priority Claimed Under 35 USC 119

I hereby claim the benefit under Title 35, United States Code § 119(e) of any United States provisional application(s) listed below:

Application No.	Filing Date

I claim the benefit under Title 35, United States Code, Section 120 of any United States application(s) listed below and, insofar as the subject matter of each of the claims of this application is not disclosed in the prior United States application in the manner provided by the first paragraph of Title 35, United States Code, Section 112, I acknowledge the duty to disclose material information as defined in Title 37, Code of Federal Regulations, Section 1.56 which occurred between the filing date of the prior application and the national or PCT international filing date of this application:

Application No.	Date of Filing	Status

**POWER OF ATTORNEY:** As a named inventor, I hereby appoint the following attorney(s) and/or agent(s) to prosecute this application and transact all business in the Patent and Trademark Office connected therewith.

Andrew T. Serafini, Reg. No. 41,303  
 William M. Smith, Reg. No. 30,223  
 Randolph T. Apple, Reg. No. 36,429

Send Correspondence to: <b>Andrew T. Serafini, Ph.D.</b> <b>TOWNSEND and TOWNSEND and CREW LLP</b> <b>Two Embarcadero Center, 8<sup>th</sup> Floor</b> <b>San Francisco, California 94111-3834</b>	Direct Telephone Calls to: (Name, Reg. No., Telephone No.) Name: Andrew T. Serafini, Ph.D. Reg. No.: 41,303 Telephone: 650-326-2400
--	---

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I further declare that all statements made herein of my own knowledge are true and that all statements made on information and belief are believed to be true; and further that these statements were made with the knowledge that willful false statements and the like so made are punishable by fine or imprisonment, or both, under Section 1001 of Title 18 of the United States Code, and that such willful false statements may jeopardize the validity of the application or any patent issuing thereon.

Signature of Inventor 1

Peter S. Lu

Date

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